

Final Technical Performance Report

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"Energy Assisted Epitaxy of GaN Using a Low Flux Nitrogen Atom Source"

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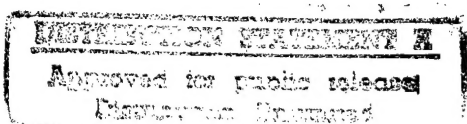
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**Final Technical Report : Energy Assisted Epitaxy of GaN Using a Low Flux
Nitrogen Atom Source**

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1.0 Program Summary

This grant was issued under the Department of Defense Experimental Program to Stimulate Competitive Research as part of a national effort to develop nationally competitive programs in states which have historically not been successful at obtaining Federal Funding for independent research. In addition, the grant provided funding to both conduct research and to educate scientists and engineers in areas important to national defense. The grant was successful in all three areas. As detailed in this report, we have performed high quality research on fundamental issues of growth in GaN, a strategic material, in addition to training many students. Finally, a nationally competitive program was established as evidenced by the award of a regular ONR grant, "An Investigation of the Effects of Hydrogen on Growth Kinetics and Defect Formation in Group III-Nitride Semiconductors", (ONR-N00014-96-1-1008) as a direct offshoot of research performed on this grant.

This report describes the research performed on the grant. Much of the research has been described in detail in the Master's Thesis of one of the students supported and in the five publications resulting as part of the program. These have been included as Appendices, and the information contained therein will thus only be summarized in the report. The areas covered in this report can be grouped as follows:

- » Basic system development and nitrogen source characterization.
- » The relationship between nucleation domains and resulting defect density and electrical quality.
- » Basic growth regimes.
- » The influence of ions.
- » The influence of uv illumination on growth kinetics.
- » The influence of atomic hydrogen on growth kinetics.
- » The role of inversion domains in determining surface morphology.
- » Student involvement.

2.0 Basic system development and nitrogen source characterization

System development and source characterization are described in detail in Appendix A, Steven Buczkowski's Thesis for his Master of Science in Physics. In summary, the GaN layers for this study were grown at West Virginia University by MBE in a custom system using a combination of an ion pump, titanium sublimation pump and cryopump. The ion and sublimation pumps ensured a clean background prior to growth while the high throughput (1000 l/s Cryotorr 8) cryopump was used during growth to adequately pump away unreacted nitrogen. A standard MBE source (EPI-40M) provided the Ga flux. For the bulk of the study, a cryogenically-cooled rf plasma source (Oxford Applied Research CARS-25) operating at either 500 or 600W was used to produce the active nitrogen flux. All layers grown with this source used a nitrogen flow rate of 6 sccm, controlled by a mass flow controller, resulting in a system background pressure of 6×10^{-5} Torr during growth. We found that our growth rates were ultimately limited to less than 1 Å/s by the maximum flux of active nitrogen available from our source configuration. The distance between our nitrogen source and the growing layer was about 30 cm, which is nearly double that found in most MBE systems. Although larger growth rates are reported for growth with the more standard source-to-substrate distances, the ultimate rate continues to be limited by the total active nitrogen flux from the plasma source used. In the latter stages of this study, we evaluated an EPI Unibulb rf source built to our specifications. More source details and characterization are summarized at the end of this section.

Atomic hydrogen was produced using a thermal cracker (EPI). Typically, 1×10^{-6} Torr beam equivalent pressure (BEP) of hydrogen was passed through the thermal source operating at 9.5 amperes, although the hydrogen flux was varied for several samples. Literature supplied with the source indicated a dissociation efficiency of about 10% for this operating condition. Therefore, the sample was exposed to both atomic and molecular hydrogen during growth. Photoluminescence (PL) measurements were performed using the 325 nm He-Cd laser line focused to a power density of 5 W/cm^2 , a grating monochromator and standard detection electronics. Hall, x-ray diffraction and

atomic force microscopy (AFM) (Digital Instruments Nanoscope II) measurements were also made.

Determination of substrate temperature and growth rate were important in this study. A spring-loaded Type K thermocouple was in intimate contact with the back of the molybdenum sample block with the substrate mounted with indium. Prior to this sequence of growths, an additional thermocouple was attached to the front of a mounted substrate and a calibration curve determined between the front and back thermocouples. This calibration was routinely checked by using the melting points of various metals, oxide desorption from GaAs, and the use of an optical pyrometer for the higher temperatures. Temperature determination was reproducible to ± 5 °C.

Growth rate was determined using total sample thickness and the growth time. Initially, the 1cm² sapphire substrates used in this study were found to frequently fall off the substrate block during growth, and so a retaining mask was used for preliminary growths in this study. While this precluded the use of our reflection high energy diffraction (RHEED) system, the masking provided a step at the samples edge allowing thickness measurement with a stylus surface profilometer (Dektak). In addition, the thickness at the sample's center was determined from interference fringes in optical transmittance measured using a Cary-14 spectrophotometer. The edge and center thickness agreed to within 10%. Later growths were performed on larger substrates that were adequately held on by indium alone. RHEED studies were performed on the growths to identify various growth regimes.

All samples were grown on c-plane sapphire substrates (Union Carbide Crystal Products). Prior to growth, the substrates were degreased and etched in a phosphoric/sulfuric (1:3) acid mixture. Initially this mixture was heated to 80°C, but a RHEED study of just the sapphire substrate alone indicated that reproducible surfaces giving a high quality RHEED pattern could only be obtained by a 10 min etch in this solution at 130 °C. Buffer layers were grown by heating the substrate to 730 °C under an atomic hydrogen flux for 20 minutes and then cooling to various temperatures for the growth of a 100 to 200 Å thick GaN buffer layer. Buffer layer growth was initiated by simultaneous exposure to the Ga and N flux. This nucleation layer was then annealed at

730 °C for 20 minutes under nitrogen flux, cooled to the growth temperature, and growth was resumed. As discussed later, these conditions represent buffer layer growth under highly Ga-rich conditions. However, after the 730 °C anneal, examination of the buffer layers by AFM indicated continuous films with no evidence of Ga condensation.

Extensive characterization of the Oxford rf plasma source was performed as described in Appendix A. We found that, although potentially orders of magnitude smaller than for an ECR source, the flux using a standard aperture contained a significant number of high energy ions. This ion content could be varied by changing the diameter of the exit holes in the aperture plate, probably due to wall interactions. Further studies were performed in the growth system with various aperture plates containing holes of differing size and number, such as to preserve the effective conductance. This data is summarized in Table 1. As can be seen, smaller hole size lead to both lower concentrations of ions as well as reduced ion energy. Unfortunately, growth rate also seemed to correlate more with aperture hole size than effective conductance. However, the ability to change the ion distribution by changing apertures allowed us to investigate the effect of relatively small amounts of ions on film growth for various energies.

After determining the low actual incorporation and "atomization" efficiency of the Oxford source, we decided to obtain a second rf source. Interaction with EPI Vacuum Products based on our characterization of the Oxford source lead to a prototype "Unibulb" source being fabricated with a large effective conductance consisting of 400 small (0.2 mm) holes. Preliminary characterization of this source is included in Table 1. A different antenna geometry apparently results in a higher "atomization" and nitrogen incorporation efficiency. Note also that direct comparison of ion content in the growth chamber (using our beam flux monitor as a Langmuir probe) indicates a lower total ion flux. Subsequent measurement supported a significantly lower ion content, with an energy distribution of the remnant ion flux comparable to the Oxford source..

Table 1. Preliminary Comparison of Oxford and EPI RF Sources

SOURCE	APERTURE		Ion Content (%)	Max. Ion Energy (eV)	Max Growth Rate ($\mu\text{m/hr}$)	Atom Efficiency (%) ^a	Incorporation Efficiency (%) ^b
	Hole Diameter (mm)	Number of Holes					
OXFORD CARS	3.0	1	5	>65	^c	^c	^c
	1.0	9	0.1	40-50	0.33	5-6	0.48
	0.5	37	0.05	15-25	0.22	3-4	0.32
	0.2	255	0.03	8-12	0.16	2-3	0.23
EPI Unibulb	0.2	400	0.01	^c	0.83	70 ^d	7.2

^a atomic N out/molecular N into source x100% [at 600W, 6 sccm, by mass spectroscopy]

^b N incorporated in GaN/molecular N into source x100% [from growth rate at 1" source-to-substrate distance]

^c not measured

^d projected from growth rate increase at 600W, 1 sccm flow

A final area of concern for the rf sources is that the nitrogen plasma apparently attacks the PBN liner as we were able to measure a significant boron content in GaN layers grown by both sources. However, GaN grown with the Oxford source contained about 10^{22} B/cm³, as compared to 10^{17} B/cm³ for the EPI source possibly due to differences in design.

3.0 The relationship between nucleation domains and resulting defect density and electrical quality

At the beginning of this study, we tried to replicate the nucleation and growth procedures listed for MBE growth of GaN at that time, which primarily used ECR plasma sources. We found that the sapphire substrate could not be nitrided using an rf plasma source, as discussed in Appendix E. In addition, the resulting layers contained a high density of very small nucleation domains. Since the defects formed during the coalescence of the nucleation domains are stable during growth, this ensured defect densities greater than 10^{12} /cm². As detailed in Appendices A, B and D, we then pursued optimization of nucleation conditions to obtain large nucleation domains, resulting in conditions which gave domains with mean diameter of 0.3 to 0.5 μ m. If there are no other defects, then this should result in defect densities of about 10^9 /cm², which we observed on several samples. The crystal quality of our material, however, continues to be limited by another pernicious defect, inversion domain boundaries as described later.

Interestingly, other groups using rf plasma sources for MBE have focused on optimizing nucleation layer growth using electrical, rather than structural measurements. This is in contrast to our work where we focused on increasing domain size first, and then began to study electrical properties. In particular, these two different approaches led to similar conditions for growth of the nucleation layer, [1] and led to electrical properties comparable to most GaN grown by MBE.

4.0 Basic growth regimes.

As described in Appendices A, B, C D and E, there are clearly several distinct growth regimes for GaN grown by rf-plasma MBE. Figure 4.1 illustrates the main points.

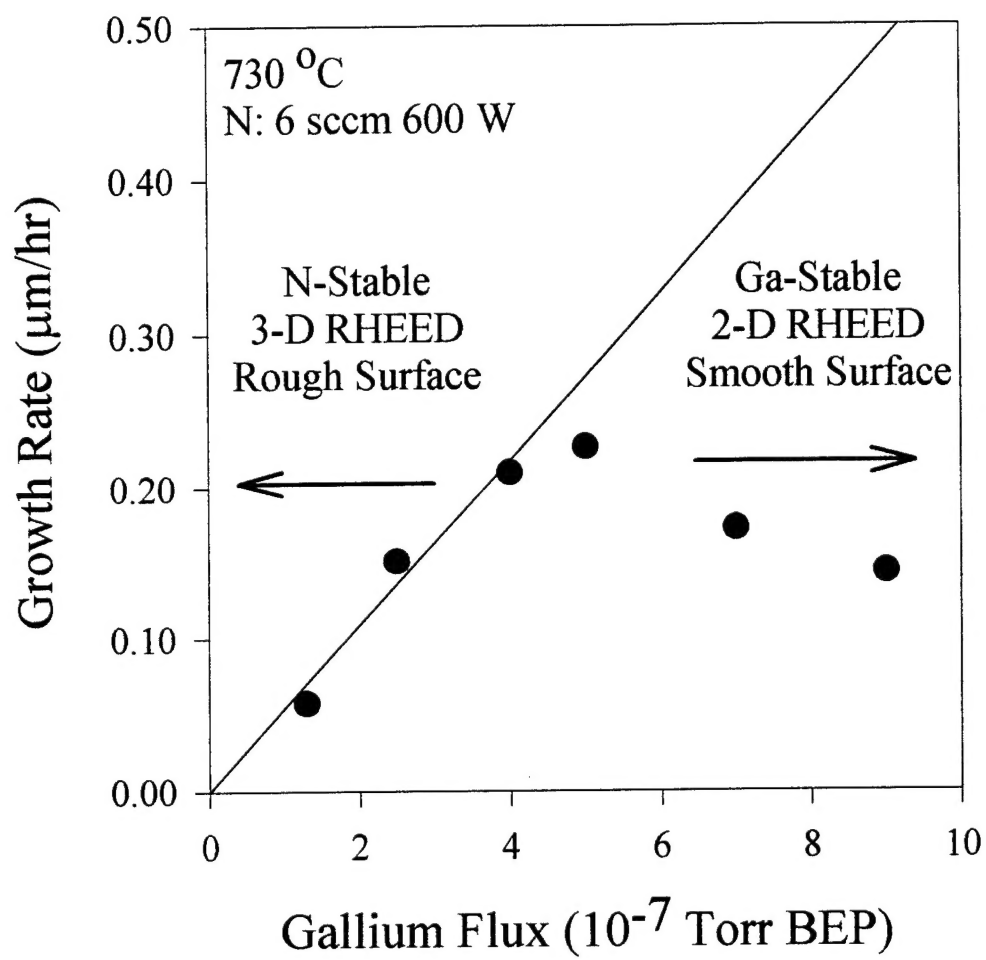


Figure 4.1 Growth rate of GaN for various conditions.

As seen in Fig. 4.1(a), increasing the Ga flux at a fixed nitrogen flux results in an increasing growth rate up to the point that the Ga is equal to the available active nitrogen. Further increases in Ga flux then give a constant or decreasing growth rate, until finally Ga condensation occurs. The former is often referred to as N-rich or N-stable growth, while the latter is Ga-rich or Ga-stable growth. As discussed in detail in the appendices, N-stable growth lead to a highly textured growth, with spotty RHEED patterns indicative of a 3-D growth mode, as shown in Fig. 4.2 (b). The majority of this surface texture consisted of pyramidal hillock-type structures related to the presence of inversion domains. The origin of these structures is more completely discussed later. In contrast, growth under Ga-stable conditions led to much smoother surfaces, with streaky RHEED patterns more indicative of a 2-D growth mode, indicated by Fig. 4.2 (a). The change-over from streaky to spotty RHEED patterns was fairly sharp, and if the 2-D growth mode was recovered fairly quickly, this was reversible. The latter allowed us to investigate the effect of external energy sources on this aspect of growth kinetics.

Figure 4.3 illustrates the other major demarcation in growth regimes. For fixed Ga and N fluxes, the growth rate is fairly constant in temperature up to some critical temperature, whereupon it begins to decrease rapidly with increasing temperature. The conventional wisdom is that this represents the onset of rapid desorption of Ga from the growing layer. Interestingly, we found that we completely eliminated the so-called "yellow-luminescence" by growing our layers above this critical temperature. Yellow photoluminescence has been attributed to various types of defects, but the actual origin is not clear. Our observation would indicate that it is associated with Ga incorporation. Further study of the temperature-dependent kinetics of basic growth is warranted, and must include some type of desorption mass spectroscopy as planned on our other ONR program.

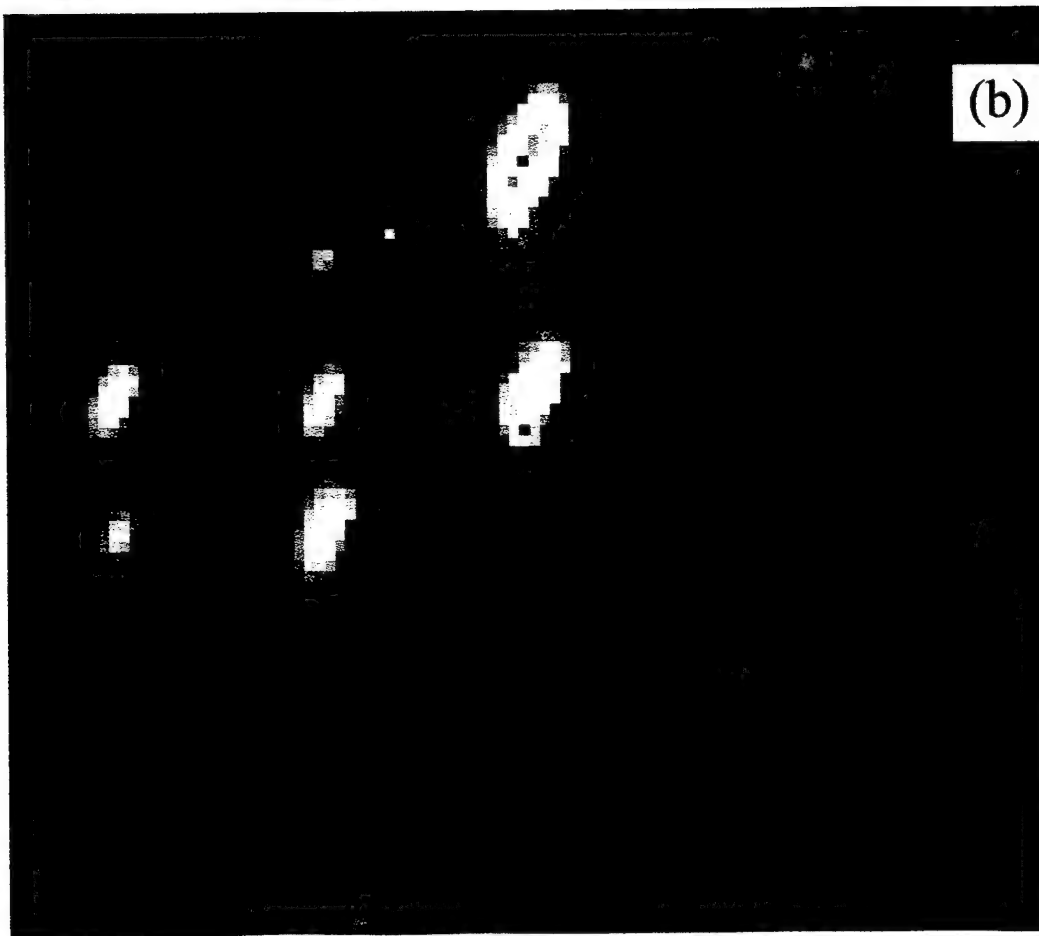
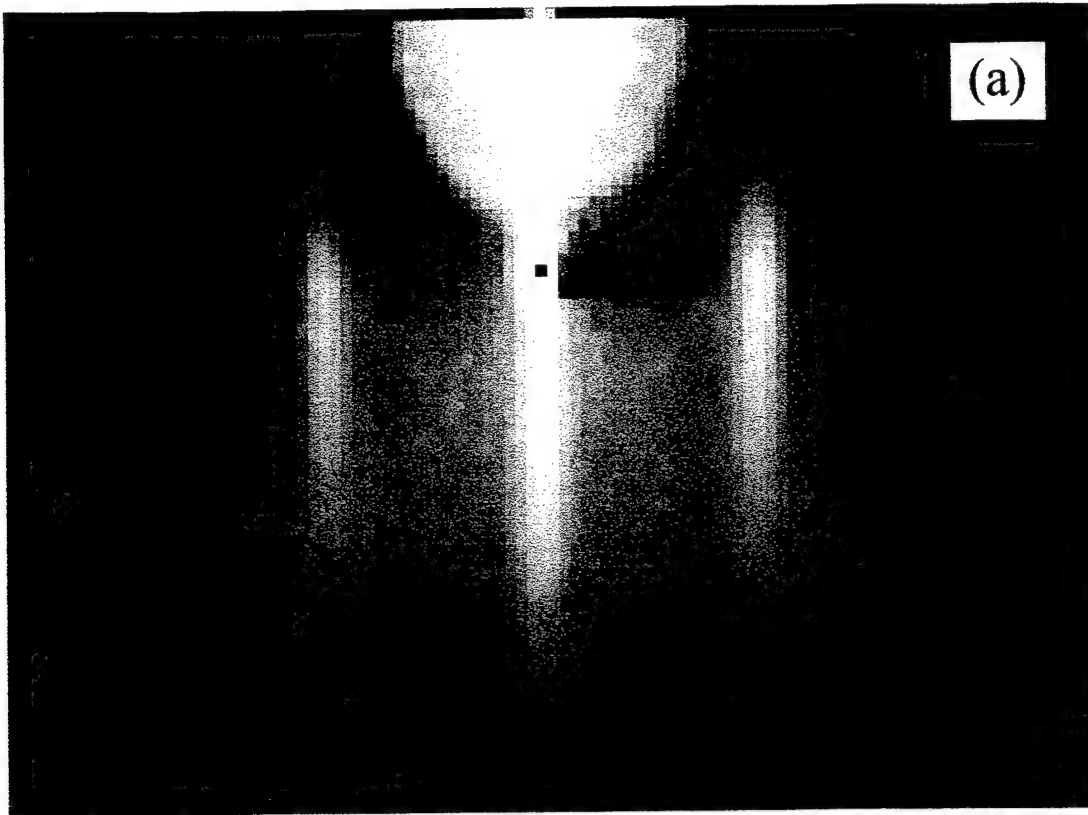


Figure 4.2 RHEED of GaN grown under (a) Ga-stable and (b) N-stable conditions.

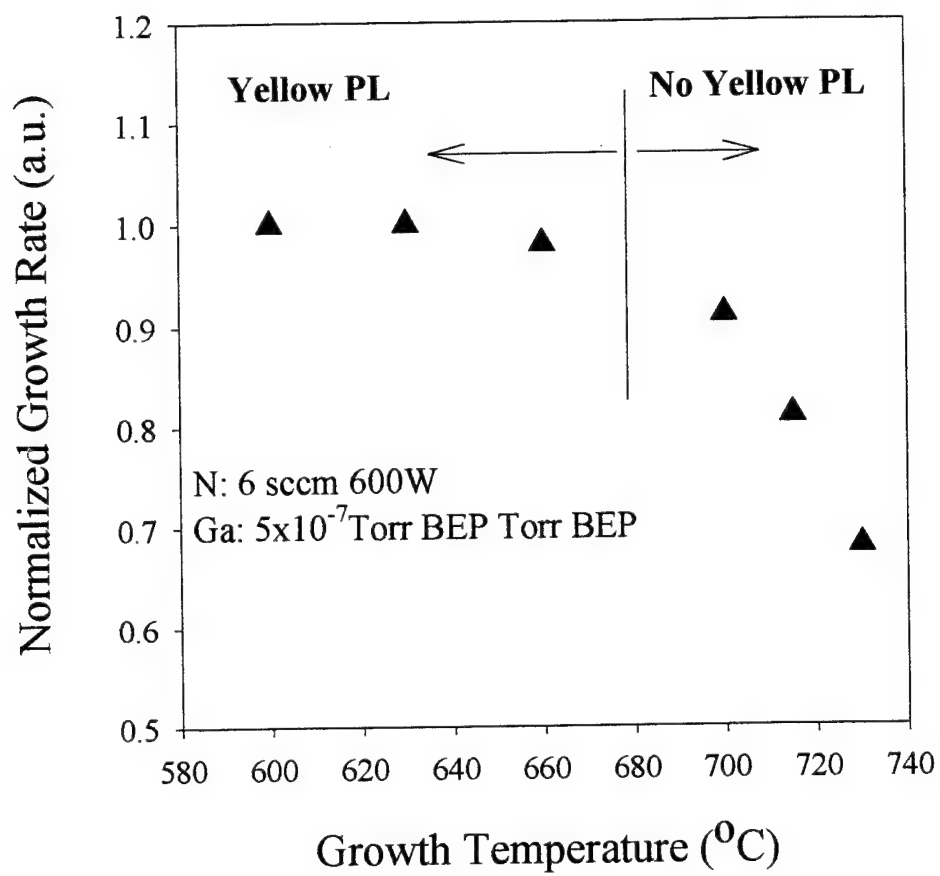


Figure 4.3 GaN growth rate as a function of temperature.

5.0 The influence of ions.

The use of different apertures on our Oxford source allowed us to qualitatively assess the effect of varying the amount of ions in the growth flux. Initially, we used the standard 37-hole aperture provided by Oxford, which we had determined had an ion content of about 0.05% with maximum energies in the range of 15-25 eV. In general, as detailed in the Appendices, we obtained material with comparable structural, optical and electrical properties to the best GaN grown by MBE. For example, typical good samples had room temperature mobilities in the range 90 to 120 cm²/Vs, and carrier concentrations of 1 to 10 x 10¹⁷ cm⁻³. (As an aside, we had one sample grown at 630 °C that had a mobility of 280 cm²/Vs and carrier concentration of 1.0 x 10¹⁷ cm⁻³ but this result could not be repeated.)

We have had private conversations with several other groups using an Oxford source, and most have been using larger apertures than the standard one as they have found they get an increase in the growth rate as we have also reported. However, this may not be the best approach. We have grown a total of 22 layers using our 9-hole aperture, which maintains the effective aperture of the normal growth aperture. From Table 1, it can be seen that the ion content approximately doubles. Perhaps of more importance, the maximum ion energy is now increased to 40-50 eV. Although we could achieve sample growth exhibiting streaky RHEED patterns indicative of 2D growth resulting in layers exhibiting photoluminescence and AFM results comparable to before, we found the electrical properties suffered significantly. While carrier concentrations were comparable at 3 to 8 10¹⁷ cm⁻³, the largest mobility was 20 cm²/Vs with more typical results around 10 cm²/Vs. **Thus we are of the opinion that ions greater than 25 eV are detrimental to growth, and should be avoided.**

We also grew several layers with the Oxford source using the 255-hole aperture. While this lowered both the ion content and the maximum ion energies, we could detect no improvement over the standard growth aperture. At this time, other issues may be masking any ion-related effects. For example, the low growth rate obtained with this aperture limited film thickness to about 0.5 µm. Based on our belief that ions must be minimized, we have worked with EPI Vacuum Products to develop an rf plasma source

based on small apertures which increased the growth rate for GaN by about a factor of eight while maintaining a low total ion content. We will be using this new source on the current ONR project, and have included a grid which can be used to suppress the residual ion content.

6.0 The influence of uv illumination on growth kinetics.

There is significant evidence that illumination during growth can shift growth kinetics. The responsible mechanism for altering the growth kinetics are, however, still poorly understood. A previous report had indicated that illumination may influence GaN growth kinetics [2] in that layers grown under illumination appeared to have a smoother surface morphology than for non-illuminated samples grown under otherwise identical conditions. Thus, we decided to investigate the influence of uv illumination on growth kinetics.

Several samples were grown under uv illumination. Comparison of samples grown with and without uv illumination under nominally Ga-rich conditions did not exhibit a significant difference in surface morphology. However, PL measurements indicated that the optical properties of the GaN may indeed be improved by the presence of uv illumination. Fig. 6.1 (a) shows the PL spectrum typical of an 0.6 μm layer grown under just Ga-rich conditions. This can be compared to the PL for samples grown under uv illumination, as shown in Fig. 6.1 (b). This latter PL is representative of thicker samples or samples grown under more Ga-rich conditions.

In an attempt to better understand the underlying mechanisms, we performed several growths where we mapped out the transition from 2-D to 3-D RHEED patterns. The results are depicted in Fig. 6.2. There is a distinct shift in the growth conditions resulting in this transition with uv illumination. One possible interpretation of this trend is that uv illumination shifts GaN growth to more Ga-stabilized conditions. This would explain the apparent improvement in surface morphology for samples grown under possibly N-stabilized (or H-stabilized) conditions [2], as well as the improvement observed in PL spectra. The underlying mechanism, however, remains unclear. For

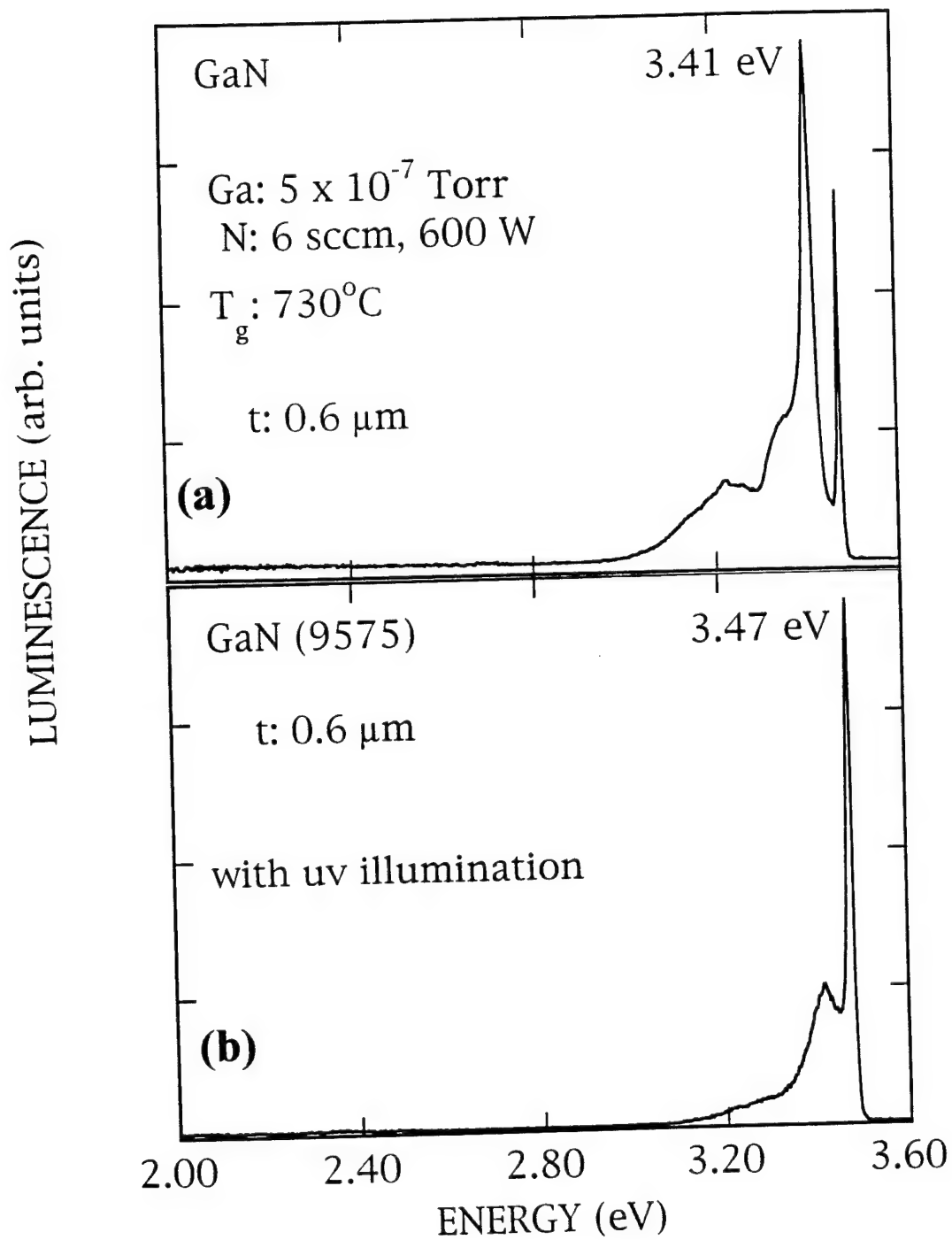


Figure 6.2 Photoluminescence of GaN grown (a) without and (b) with uv illumination.

Influence of uv Illumination on Growth Kinetics

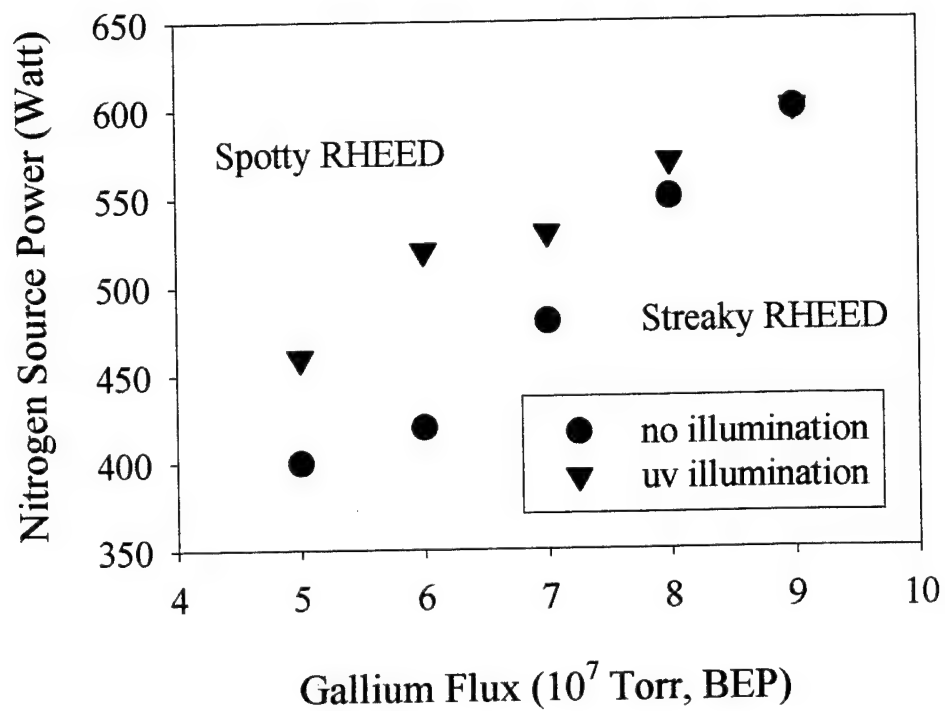


Figure 6.2 Transition from "spotty" to "streaky" RHEED patterns with and without uv illumination.

example, lowering the sticking coefficient of N or increasing that of Ga may result in the same effect. We note in passing that there was not any apparent effect on growth rate. Thus, we observe a distinct effect with uv illumination, but feel that additional experiments involving desorption mass spectroscopy are required to more clearly understand the effect on growth kinetics. We hope to include such experiments as part of our current ONR contract, "An Investigation of the Effects of Hydrogen on Growth Kinetics and Defect Formation in Group III-Nitride Semiconductors", (ONR-N00014-96-1-1008).

7.0 The influence of atomic hydrogen on growth kinetics.

The presence of an atomic hydrogen flux can have a dramatic influence on the growth kinetics of GaN. The effects are discussed in detail in the attached appendices, and will only be summarized here. In particular, two main effects are observed. The first is that the presence of atomic hydrogen shifts Ga-stabilized growth to what are apparently N-stabilized conditions. This shift is indicated by the combination of a shift to 3D RHEED patterns during growth, an increase in growth rate to Ga-arrival limited rates (as shown in Fig. 7.1), and the shifting to a surface morphology dominated by pyramidal hillocks. The other major effect is the apparent stabilization of the growing GaN surface to either thermal decomposition or diminished Ga-sticking coefficient, as shown in Fig. 7.2. While much more dramatic than the changes in growth kinetics due to uv illumination, the underlying mechanisms remain unclear. In particular, one needs to directly measure the effect on the sticking coefficients and desorption rates of the constituent species. Understanding the influence of atomic hydrogen on growth kinetics is one of the major goals to be accomplished using desorption mass spectroscopy on our current ONR contract "An Investigation of the Effects of Hydrogen on Growth Kinetics and Defect Formation in Group III-Nitride Semiconductors", (ONR-N00014-96-1-1008).

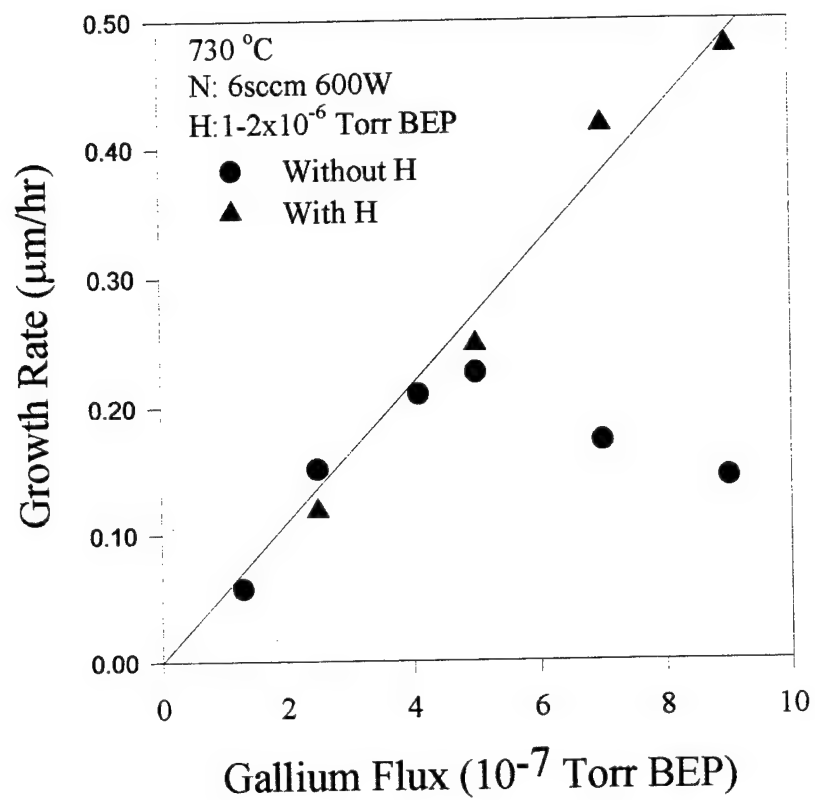


Figure 7.1 Effect of atomic hydrogen on growth rate.

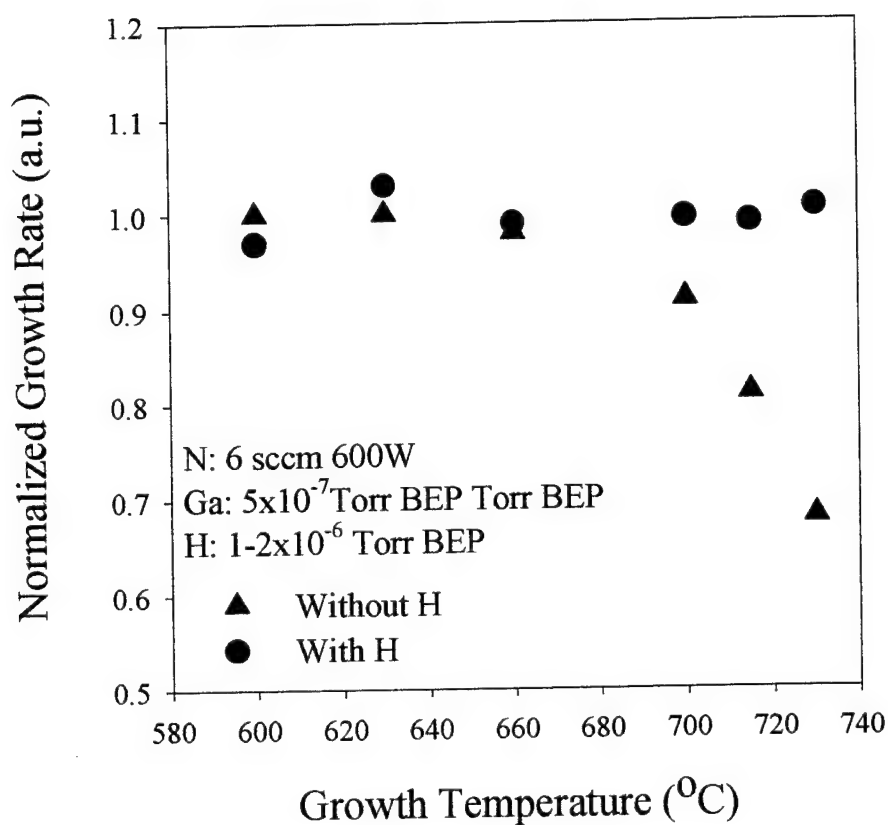


Figure 7.2 Effect of atomic hydrogen on growth rate at higher temperatures.

8. Inversion Domains and Surface Morphology

Samples grown either under N-stable conditions or under an atomic hydrogen flux often exhibit a highly textured surface morphology consisting of pyramidal hillocks. Figure 8.1 is an AFM micrograph illustrative of such a surface completely dominated by the triangular hillocks, while Figure 8.2 is from a sample exhibiting isolated hillocks. The hillocks have a triangular cross-section that range from 0.2 to 0.5 μm on a side for 1 μm thick samples, and between 150-250 nm in height. Increasing the Ga/N ratio (without atomic H) results in smaller size pyramids. These features were not observed for highly Ga-rich conditions.

Figure 8.3 shows a XTEM image of a hillock from a 1.0 μm thick sample grown under an atomic hydrogen flux with a surface that contained a high density of pyramid-shaped hillocks. The hillocks were ~ 100 nm high and ~ 200 nm wide at the base and each contained an ID with a cross-section of ~ 10 nm as shown in Fig. 8.3. The IDs were found to originate at the film/substrate interface and extend to the film surface with a constant cross sectional area between 5-20 nm. The IDBs were along the $\{10\bar{1}0\}$ planes, similar to IDs found in other films grown by MBE [3,4] and MOCVD [5,6]. What is amazing is the pronounced effect the small ID has on surface morphology. In terms of sample volume, the ID density in the sample shown in Figure 8.1 is less than 5%, yet effects associated with the IDs totally dominate the sample's morphology.

GaN layers grown by MOCVD with a similar surface morphology have been investigated by Daudin *et al.* [5] using ion channeling and convergent beam electron diffraction. They found that the IDs associated with pyramids at the surface of their samples were oriented (0001) (or Ga- terminated), while the bulk of the matrix was oriented (000 $\bar{1}$) (or N-terminated). In contrast, they were also able to grow "flat" samples which were single phase (0001) and contained no IDs. Hillock formation apparently results from the higher growth rate of the ID located at its midpoint. The resultant strain at the boundary may also enhance the growth rate of the opposite phase and dictate the final surface morphology. Perhaps this non-planar growth region is a source of point defects which degrade the properties of samples grown under N-stable conditions. [1].

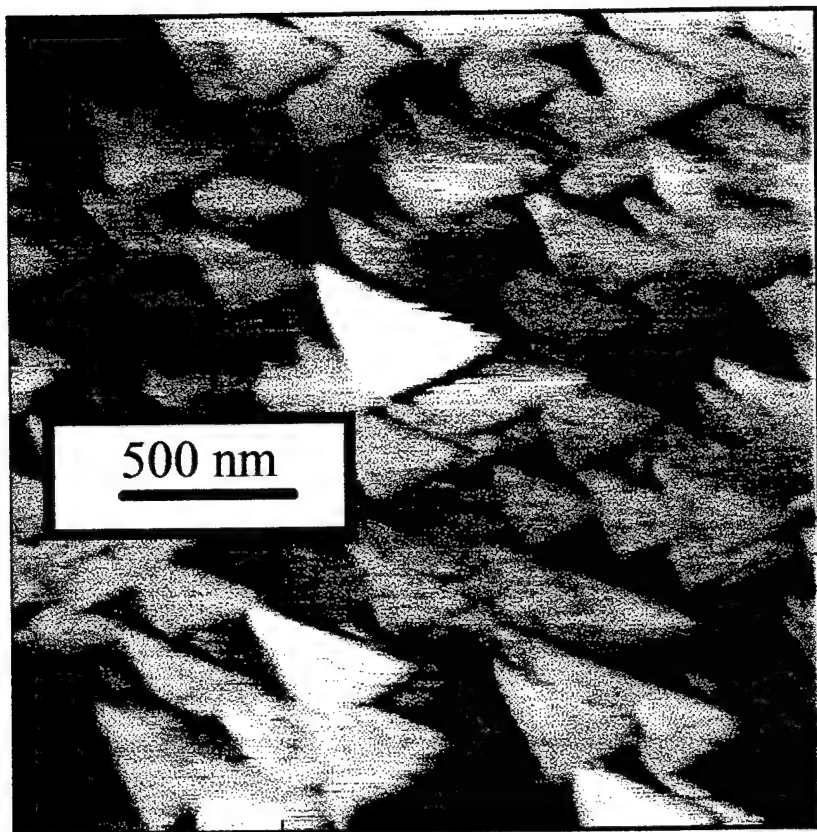


Figure 8.1 AFM of a highly textured GaN sample.

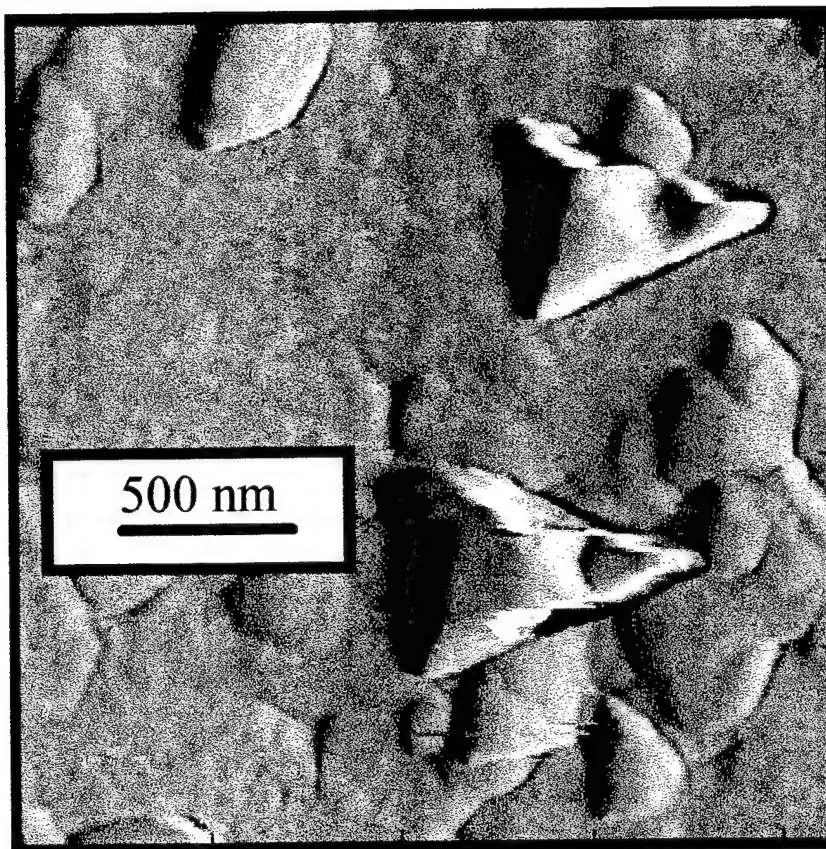


Figure 8.2 AFM showing isolated hillocks.

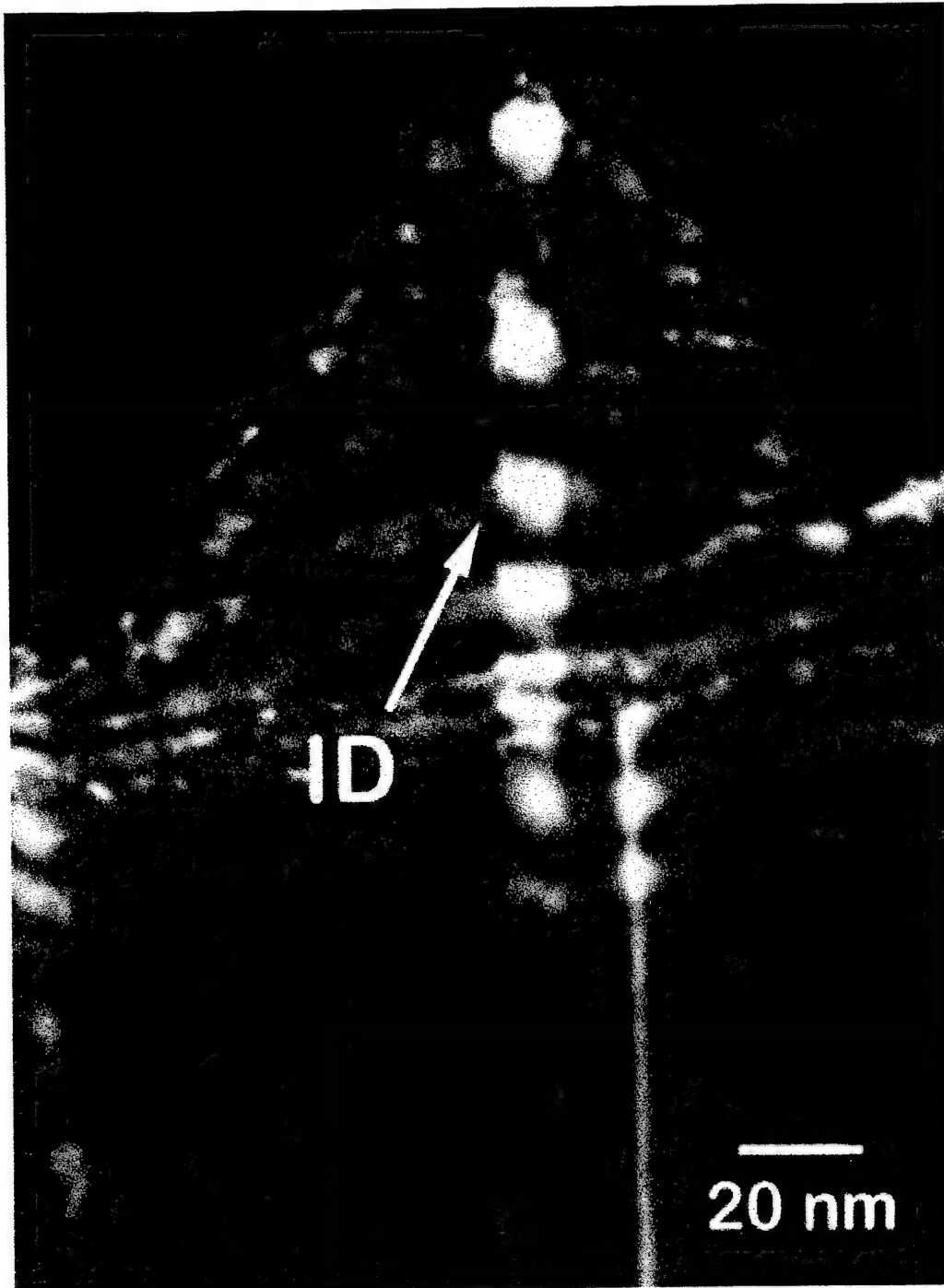


Figure 8.3 Transmission electron micrograph of a pyramidal feature showing the central inversion domain.

We believe our IDs are oriented (0001) similar to the MOCVD results [5]. This is supported by a recent study by Smith *et al.* [7] who found by STM measurements for films grown by rf-plasma MBE using similar techniques that the polarity of the matrix was N-terminated. If our assessment is correct, however, the Ga-terminated surface can have a significantly higher growth rate than a N-terminated surface, in agreement with the speculations of Middleton *et al.* [8]. The growth rate differential is also consistent with recent results on MOCVD GaN reported by Liliental-Weber *et al.*, [9] where (0001)-oriented IDs grow at a slower rate than the (0001)-oriented matrix, leading to oriented "pits" that could be described as inverted hillocks. One implication is that if the bulk matrix is nucleated to be Ga-terminated and a N-terminated ID is of small cross-section, the possibility exists for overgrowth of the ID under N-stable conditions, removing this defect after growth of a sufficient layer thickness. Thus, the observation that inversion domains are stable during MBE growth of GaN may be due to the relative differential growth rate on (0001)-oriented material.

Samples grown under excess Ga were found to be free of pyramidal hillocks, even for films containing high densities of IDBs. Again, IDBs were observed to originate at the film/substrate interface and propagate through the growing layer. In sharp contrast to N-stable growth, no significant hillock features were observed at the intersection of the IDB with the surface of the GaN. However, as reported earlier [10] the surface of the ID is found to be 2 nm higher than the surrounding matrix, indicating a slight enhancement in the growth rate in the region near the ID. This height difference is significantly less than in the film shown in Fig. 8.3. Thus, Ga-stable growth appears to suppress the growth-rate differential between the two different polarities of GaN, directly leading to the observed smoother surface morphologies.

The formation of IDBs is not an intrinsic property of MBE growth of GaN, but is apparently related to nucleation conditions. We have grown several samples under N-stable conditions that only exhibit the granular texture associated with N-stable growth, without the presence of hillocks. The IDs initiate within the buffer layer and are localized laterally with a relatively small cross-section, indicating a localized nucleation site. We obtain different densities of IDBs with what we believe are identical nucleation conditions.

This could be due the presence of steps in the sapphire surface as suggested previously [11], defects in the substrate surface itself from remnant polishing damage, or possibly related to high energy ions present in the nitrogen flux itself. (With respect to the latter, we have measured a small but finite flux of nitrogen ions with energies > 25 eV from our nitrogen source)

9.0 Student Involvement.

While supporting several students, many additional students have been influenced by this program. The following represents a list of students who were either supported by this program, co-author on a presentation or publication resulting from research related to this program, or both.

Graduate Students

Zhonghai Yu (Ph. D. (Physics))
Steven Buczkowski (M.S. (Physics))
Matt Millecchia (M.S. (Physics))
Sumeet Kumar (M.S. (Chemical Eng.))
Lauren Hirsch (M.S. (Physics), Ph. D Candidate)
Aaron Ptak (Ph. D Candidate (Physics))
Monica Moldovan (Ph. D Candidate (Physics))
Katie Ziemer (Ph. D Candidate (Chemical Eng.))
Hongtao Shi (Ph. D Candidate (Physics))
Tim Charlton (Ph. D Candidate (Physics))
Steve Frazier (Ph. D Candidate (Electrical Engineering))

Undergraduate Students

Gabriel Gobeli (Physics)
Elizabeth Mayo (Physics)
Tim Charlton (Physics)

10.0 References

1. H. Reichert, R. Averbeck, A. Graber, M. Schienle, U. Straub, and H. Tews, *Mater. Res. Soc. Symp. Proc.* **449**, 149 (1997)
2. M.J. Paisley and R.F. Davis, *J. Vac. Sci. Technol. A* **11**, 18 (1993).
3. L.T. Romano, B. S. Krusor, R. Singh, and T. D. Moustakas, *J. Electron. Mater.* **26**, 285 (1997)
4. Zhonghai Yu, S.L. Buczkowski, N.C. Giles, T.H. Myers and M.R. Richards-Babb, *Appl. Phys. Lett.* **69**, 2731 (1996); S.L. Buczkowski, Zhonghai Yu, M.R. Richards-Babb, N.C. Giles, T.H. Myers and L.T. Romano, *Mat. Res. Soc. Symp.* **449**, 197 (1997).
5. B. Daudin, J. L. Rouviere, and M. Arley, *Appl. Phys. Lett.* **69**, 2480 (1996)
6. Z. Liliental-Weber, H. Sohn, N. Newman, and J. Washburn, *J. Vac. Sci. Technol. B* **13**, 1578 (1995)
7. A. R. Smith, R. M. Feenstra, D. W. Greve, J. Neugebauer, and J. E. Northrup, *Phys. Rev. Lett.* **79**, 3934 (1997).
8. P.G. Middleton, C. Trager-Cowan, A. Mohammed, K.P. O'Donnell, W. Van Der Stricht, I. Moerman and P. Demeeser, *Mat. Res. Symp. Proc.* **449**, 471 (1997)
9. Z. Liliental-Weber, Y. Chen, S. Ruvimov and J. Washburn, *Phys. Rev. Lett.* **79**, 2835 (1997)
10. L.T. Romano and T.H. Myers, *Appl. Phys. Lett.* **71**, 3486 (1997).
11. L.T. Romano, J. E. Northrup, and M. A. O'Keefe, *Appl. Phys. Lett.* **69**, 2394 (1996)

Nucleation and Growth of GaN on Sapphire by MBE

Thesis

Submitted to the Eberly College of Arts and Sciences of
West Virginia University
In Partial Fulfillment of the Requirements for
The Degree of Master of Science

by
Steven Buczkowski
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GaN was grown by molecular beam epitaxy in an effort to determine nucleation and growth conditions which lead to high quality, single-crystal films. Atomic force microscopy (AFM) was used to show that growth under Ga-rich conditions promotes the nucleation of films with large nucleation domains leading to a lower density of intrinsic defects related to domain boundaries. These conditions are also shown to promote a 2-D growth mode resulting in films with a high degree of nucleation domain coalescence and surface roughnesses below 2 nm. Addition of atomic hydrogen, using a thermally-cracked source, is shown to increase the growth rate of Ga-rich growths by a factor of two. The hydrogen appears to be altering the growth kinetics to increase the residence time of nitrogen atoms at the substrate surface. The addition of hydrogen seems to have no effect on the material characteristics as probed by Hall measurements and photoluminescence. In addition, the RF source used to generate active nitrogen species was characterized by means of mass spectroscopy and optical emission spectroscopy. Under normal operating conditions for GaN growth, the source was shown to convert about 5% of the total nitrogen input into neutral, nitrogen atoms and about 0.03% into nitrogen ions which may cause crystal damage during growth.

This work may have never reached this state were it not for my wife Debra's constant encouragement and ability to make sense out of MS Word. For her love and support, I am truly indebted.

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1. Introduction

1.1 Wide bandgap semiconductors

Semiconductor technology has become increasingly important in the last fifty years, first with the advent of computer technology and more recently with the increasing drive toward optoelectronics. The need for smaller, faster and more robust electronic devices has created a tremendous push in the materials sciences to produce materials better suited to the tasks at hand. Within the last thirty years, optoelectronics have come to the forefront of technology as one solution to the size and speed demands imposed on modern electronics. The III-V materials GaAs and AlGaAs have been very successful in this arena. GaAs-based high frequency switching circuits are becoming increasingly available and AlGaAs based laser diodes have spawned a tremendous revolution in the area of optical storage.

Currently, semiconductor optical devices routinely operate from the infra-red (IR) to green wavelengths. By developing high reliability devices which would operate into the blue and ultra-violet (UV), semiconductor based devices could emit and detect over the entire visible spectrum impacting heavily on imaging and graphics applications. For example, a blue or UV light laser diode (LD) would increase currently diffraction limited near-IR optical storage densities by more than a factor of four.

The longest wavelength of light which a semiconductor can generate or respond to is dictated, to first order, by its bandgap energy as $E_{\text{Bandgap}} = hc/\lambda$. The larger the bandgap, the shorter the wavelength of light that can be captured or created. In order to reach the blue end of the visual spectrum then, it is necessary to develop systems with bandgaps on the order of 2 eV or higher. The best contenders in this field to date are the III-V nitride materials (GaN 3.4 eV; AlN 6.2 eV; InN 1.9 eV), ZnSe (2.67 eV) and SiC (2.2 eV). SiC has been studied very extensively and has been a major force in the blue LED market for some time. It is however limited by an indirect bandgap which reduces

its overall light emission efficiency thereby eliminating all hope of building a SiC laser diode structure. SiC does have the advantage that it is currently one of the most promising wide bandgap materials for non-optical devices, such as bipolar and field-effect transistors [1]. The II-VI material ZnSe has long been considered the next step in laser diode technology but it is still very limited by problems with high contact resistance, low p-type doping and short device lifetimes related to defects in the crystal structure [2].

The III-V nitrides may prove to be the best of all possible choices. GaN, InN and AlN form a continuous alloy system with bandgaps ranging from 1.9 eV to 6.2 eV. This would allow potential device operation ranging from red wavelengths to the far UV. The III-V nitrides are expected to have superior radiation hardness which, combined with a resistance to both high temperatures and most acids, makes them very attractive for use in space-borne electronics and harsh environments.

Considerable work has been done recently toward realizing the potential of these materials. However, even with the rapid advancement in material quality and the exhibition of numerous practical devices, many issues have yet to be resolved fully. Most groups have focused on molecular beam epitaxy (MBE), chemical vapor deposition (CVD) or their variants as the growth system of choice [3]. While metal organic CVD (MOCVD) offers significantly higher growth rates, making it attractive for commercial device applications, the presence of the large amounts of hydrogen typical of MOCVD has been shown to heavily compensate p-type dopants [4,5]. This has been overcome somewhat by the finding that dopant passivation can be partially reversed by either low energy (10-15 keV) electron beam irradiation or a thermal anneal in a N₂ atmosphere [6,7]. These findings have recently led to the commercial availability of blue LEDs, based on GaN, by Nichia Chemical Industries in Japan [8] and CREE Research, Inc. in the United States. MBE, on the other hand, offers an environment free of hydrogen for p-type material growth with no need for post-growth treatment. However, it is hindered by

significantly lower growth rates and materials properties which are generally inferior to MOCVD grown material.

CVD related growth typically uses ammonia to provide a flux of atomic nitrogen. At the higher substrate temperatures (1050°C) involved, the ammonia molecule is readily decomposed to provide nitrogen at the growth surface. In order to maintain the relatively hydrogen free MBE environment, most MBE researchers have turned to using nitrogen gas. Due to the high bond strength (9.5 eV) of the N-N bond, thermal dissociation at the substrate, alone, produces an intolerably low flux of active nitrogen. To overcome this, MBE growth of GaN based on molecular nitrogen requires an auxiliary method for breaking the N-N bond. MBE researchers have split among four principle methods for overcoming this energy barrier. Most have turned to one of two types of plasma sources, either radio frequency (RF) or electron cyclotron resonance (ECR), for this bond breaking. Others have utilized either ion sources or ammonia to provide reactive nitrogen. For the plasma and ion sources, questions still remain as to the identity of the species responsible for film growth. Both neutral nitrogen atoms and low energy ions have been shown to be beneficial while high energy ions have been shown to have deleterious effects on sample quality[9,10]. Ammonia has been used successfully in a technique commonly referred to as gas source (GS)MBE to combine aspects of MBE and MOCVD. Researchers using this technique have demonstrated MOCVD-like growth rates but questions still remain as to the effects of introducing a large hydrogen flux into the growth[11].

All Group III-nitride growths, regardless of process, share two common problems. Large single crystal substrates have yet to appear for the III-nitrides so all growth must be heteroepitaxial, and samples grown without intentional dopants still exhibit a high degree of n-type conductivity.

Heteroepitaxial growth itself is not a fundamental problem. However, for GaN in particular, no sufficiently lattice matched substrate has been found to date (Figure 1.1).

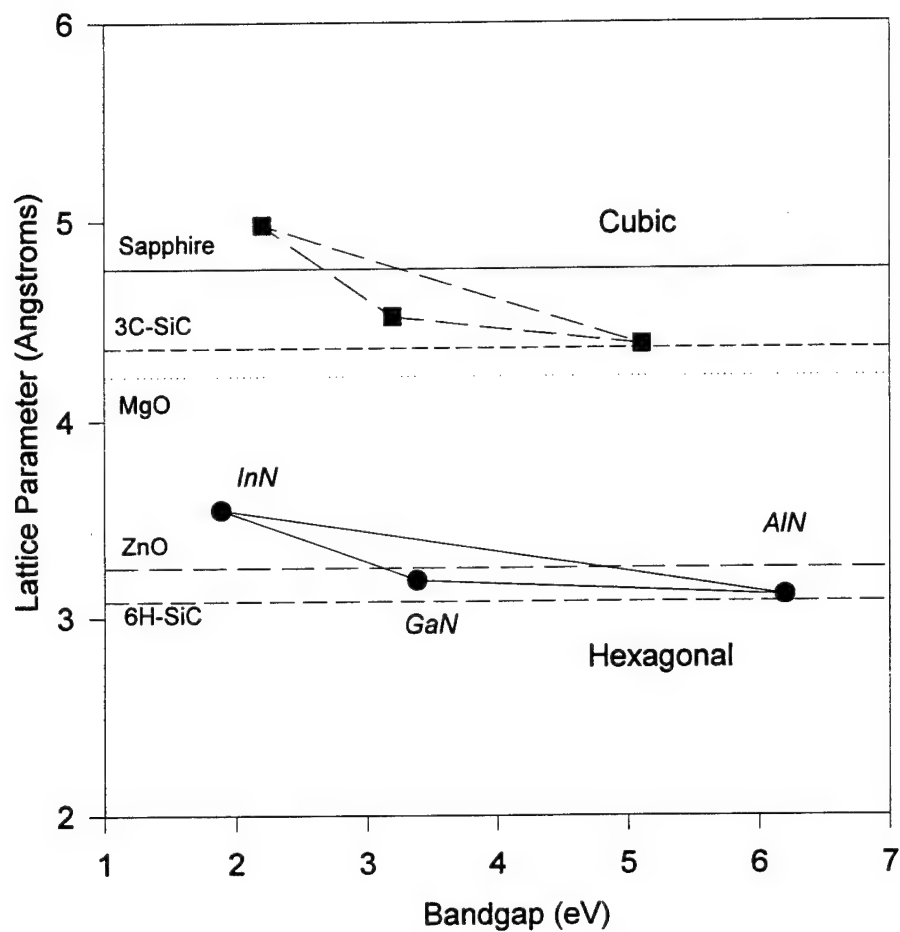


Figure 1.1 Schematic comparing lattice matches for the III-V nitrides and several common substrates. The system of interest here is the sapphire to hexagonal GaN. The hexagonal values are for the in-plane, or 'a', spacing.

Attempts have been made to grow GaN on sapphire ($\alpha\text{-Al}_2\text{O}_3$), ZnO, MgO, SiC, GaAs, and spinel (MgAl_2O_4), among others. Even though it is one of the least lattice matched substrates available, sapphire is the most common substrate for GaN growth due to its availability and the convenience of pre-growth handling. Sapphire's popularity is also attributable to the successful use of buffer layers grown between the substrate and the epilayer to improve the morphology and electrical character of films grown in severely mis-matched systems. In spite of the lattice mismatch, working LED structures are now commonly grown on sapphire substrates. Due to their better lattice match, ZnO, MgO, and SiC are expected to become more popular as they become more readily available.

GaN, in particular, has a large problem with unintentional n-type auto doping. Undoped samples typically exhibit carrier concentrations in the 10^{17} to 10^{18} cm^{-3} range. This has been attributed to either nitrogen vacancies or oxygen impurities[12]. With the recent improvements in growth practices, films are now commonly grown with carrier concentrations as low as $5 \times 10^{16} \text{ cm}^{-3}$ and mobilities up to $1000 \text{ cm}^2/\text{Vs}$ at room temperature[13].

Finally, the chemical resistance of the wide bandgap materials has made it difficult to etch device structures in the grown material. This however has been mostly overcome by the use of plasma/ion beam assisted etching. A comprehensive discussion of wide-bandgap semiconductors can be found in the review articles by Strite and Morkoç[1,14].

1.2 Motivation and Goals of this Research

The current research program in GaN at WVU is directed at understanding the processes involved in the growth of GaN, and includes investigating how these processes may be affected by the addition of non-growth species, such as hydrogen, or non-thermal energy such as irradiation with above bandgap light or high energy electrons. The work reported in this thesis represents the end-product of an effort to establish a baseline

understanding of GaN growth and, as such, has defined directions for more detailed investigations.

The specific goals of this research project were:

1. Identify the various active nitrogen species produced by a RF nitrogen source for a variety of source operating parameters, such as, pressure, gas flow rate and plasma power.
2. Develop an understanding of the buffer layer nucleation, particularly to control the initial domain size and crystal polarity.
3. Establish a baseline for GaN growth, with particular emphasis on determining conditions which promote two-dimensional growth modes.
4. Perform a preliminary investigation of the effect of hydrogen on GaN growth kinetics through use of a thermal cracker to inject a flux of atomic hydrogen during growth.

2. Background

2.1 Summary of GaN material characteristics

The following table lists some of the accepted values for the material characteristics of GaN in its two most common polytypes. The wurtzite phase is the most common of the tetrahedrally-bonded polytypes seen in GaN and results from growth on a hexagonal substrate. The zincblende polytype is the next most common, resulting from growth on a cubic substrate, and appears to have a slightly higher energy of formation.

Fundamental properties of GaN

	<i>Wurtzite Polytype:</i>	<i>Zincblende Polytype:</i>
Band-gap Energy	$E_g(300K) = 3.39\text{eV}$ $E_g(1.6K) = 3.50\text{eV}$	$E_g(300K) = 3.2\text{-}3.3\text{eV}$
Lattice constants	$a = 3.189\text{\AA}$ $c = 5.185\text{\AA}$	$a = 4.52\text{\AA}$
Coefficients of Thermal Expansion	$\Delta a/a = 5.59 \times 10^{-6} \text{ K}$ $\Delta c/c = 3.17 \times 10^{-6} \text{ K}$	
Thermal conductivity	$\kappa = 1.3 \text{ W/cm K}$	
Index of refraction	$n(1\text{eV}) = 2.33$ $n(3.38\text{eV}) = 2.67$	$n(3\text{eV}) = 2.9$
Dielectric constant	$\epsilon_0 = 9$ $\epsilon_\infty = 5.35$	
Melting Point	Sublimes in vacuum at temperatures $> 800\text{-}850^\circ\text{C}$	

2.2 GaN growth on Sapphire

Since GaN single crystal substrates do not exist at this time, growth must be performed heteroepitaxially using substrates based on other material systems. For GaN this has included sapphire (α -Al₂O₃), ZnO, MgO, Si, SiC, GaAs, Spinel (MgAl₂O₄) and others. Even though ZnO, MgO, and SiC are more closely lattice-matched to GaN, sapphire has been the substrate of choice for many research groups because of its availability and ease of preparation prior to growth. Growth on sapphire has been attempted on a wide variety of crystal orientations with the (0001), ($\bar{1}012$) and (10 $\bar{1}0$) orientations showing the best growth relationships with GaN [15].

Despite its popularity, sapphire presents a major difficulty to the growth of GaN. Its large lattice mismatch, $\delta = [(a_{\text{GaN}} - a_{\text{Sapphire}}) / a_{\text{Sapphire}}]$, of 33% favors the formation of small, isolated three-dimensional growth islands which do not easily coalesce into large-scale single crystal growth. The boundaries of the islands typically form edge-type dislocations which are difficult to eliminate in subsequent growth. These defects then affect the electrical and optical characteristics of the material by acting as traps or scatterers. One of the main thrusts of GaN work in general, and this work in particular, has been to optimize the growth conditions for GaN on sapphire to overcome these mismatch-induced dislocations. This issue can be approached on two fronts by finding growth conditions favorable to the elimination of existing dislocations (2-D growth), and also by promoting nucleation of larger islands so that there are fewer domain edges to generate dislocations from the start. The present work focuses more on the latter of these fronts with an ultimate goal being conditions which promote both.

The most common method for relieving the mismatch before GaN growth has been to grow a buffer layer with an intermediate material. Originally, most groups used a thin (200-400Å) AlN layer grown at a low temperature (500°C) and subsequent GaN

growth at a higher temperature (800-1000°C). The low temperature AlN layer was shown to be a uniform, amorphous layer on the sapphire which then re-crystallizes during the high temperature GaN growth. This approach seems to accommodate the strain between the three material layers. This same technique has since been tried with a GaN buffer layer with equal success. Kuznia *et al.*, [16] performed MOCVD growths to determine which material made the better buffer layer for sapphire. Their findings, by electron and x-ray diffraction, indicate that both AlN and GaN are equally suited and either is significantly better than no buffer layer at all. X-ray data showed a factor of three improvement in rocking curve FWHM with the buffer layers. They also discovered that the buffer layers had an optimal thickness which promoted the highest crystallinity and the best electrical characteristics in the actual epilayer.

In addition to buffer layers, most groups expose the sapphire to a flux of active nitrogen prior to growth. The belief is that this surface nitridization produces a layer of AlN which becomes a part of the buffer layer system.

2.3 MBE Growth Kinetics

Before discussing how changing growth parameters affects film growth, some groundwork needs to be laid in the process of film growth itself. One of the goals in semiconductor growth is the growth of atomically smooth, single-crystal epitaxial thin films. This means that the grown film is crystallographically related to the substrate, with all of the film sharing this relationship. One of the benefits of such growth is that nearly atomically smooth surfaces allow the sharpest intermaterial junctions making such things as quantum well structures possible. Electrically and optically, a single crystal material offers the least resistance to the passage of charge carriers and photons.

Molecular beam epitaxy is an ultra-high vacuum (UHV) crystal growth technique which has become popular in the last twenty years. MBE derives its name from the fact

that growth occurs with material fluxes low enough that chemical interactions in the gas phase are negligible. This is in contrast to chemical vapor deposition growth, where gas phase chemistry plays a major role in the deposition process. The UHV environment also considerably reduces background contamination levels.

Figure 2.1 shows schematically the competing processes involved in the epitaxial growth of a material. Starting with an incident flux of molecules coming from an overpressure of a gas or a molecular beam from a knudsen source, some fraction of the flux adsorbs onto the substrate. Once on the substrate, adsorbed molecules can diffuse across the substrate surface until they either bond with another entity or desorb from the surface. Bonding can occur to other adsorbed molecules, which may also be diffusing across the surface, or to either a step edge formed by crystalline defects, a natural terrace in the substrate or the boundary of a nucleating cluster of material. In this kinetically-driven growth, one or more of the rates associated with these processes can become the limiting factor in the rate of growth.

Once nucleation begins, crystal growth follows one of three general paths: Frank-Van der Merwe (layer-by-layer) growth, Stranski-Krastanov (layer-plus-island) growth, and Volmer-Weber (island) growth. In the island growth mode, the impinging atoms are more strongly bound to one another than they are to the substrate. As a result, small clusters nucleate on the substrate and proceed to grow three-dimensionally, thus retaining their separate character. The layer-by-layer growth mode is exactly opposite to the island growth. Atoms are more strongly bound to the substrate, so nucleation occurs a monolayer at a time. As further layers are added, they become slightly less energetically bound than the previous layer. If the decrease in binding energy between layers decreases monotonically toward the bulk crystal value, then the layer growth continues. Intermediate to these is the layer-plus-island growth, where the initial layer-by-layer growth is disturbed through strain or dislocation formation, and three-dimensional growth ensues.

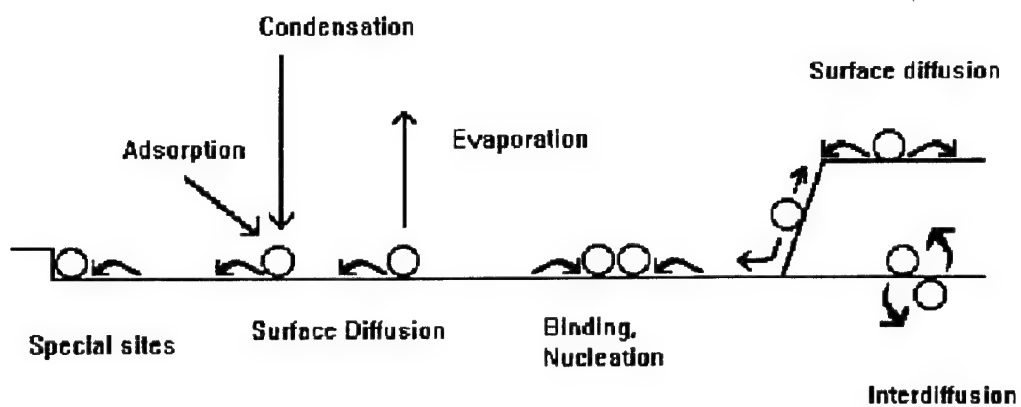


Figure 2.1 Schematic diagram of processes involved in film nucleation and growth.

In general, the 3D growth modes produce inferior materials because the inter-crystal boundaries act as traps or scatterers for charge and photons. The 2D layer growth mode is favorable since it produces the smoothest film interfaces. Also, as defects form they are pushed transversely to the growth axis by the crystal growth. By propagating 'sideways', the defects will eventually encounter either the edge of the crystal or another defect, with which it can form a closed loop defect. Either of these events terminates the defect's propagation in the crystal.

In theory, homoepitaxial growth on a clean surface should always proceed by 2-D layer growth. In practice, this is very nearly achievable. Heteroepitaxy, on the other hand, generally proceeds by some form of island growth due to lattice-mismatch or thermally induced strain. It is possible, however, to coerce a more 2-D growth pattern by careful management of growth parameters.

A more in-depth description of these processes can be found in review articles by Cho and Arthur [17], Venables, *et al.* [18], and Joyce [19].

3. Experimental

3.1 MBE System

3.1.1 Chamber Description

The Molecular Beam Epitaxy system used for the growths described in this work was designed and built in the West Virginia University Physics Department. Housed in a nominally Class-1000 cleanroom facility, the ultrahigh vacuum (UHV) system consists of two chambers which can be isolated from one another by a gate valve (Figure 3.1). The smaller of the chambers functions as a load-lock which can be rapidly cycled between atmospheric pressure and $\sim 10^{-7}$ Torr to allow sample introduction into the growth chamber with minimal disturbance to its UHV environment. The load-lock consists primarily of a stainless steel six-way cross and bellows. A four-way cross and tee attached to the top of the load-lock provide space for an ion gauge to monitor the load-lock pressure and for preliminary bake-out of source ovens before they are put into the growth chamber. A magnetically coupled linear motion feedthrough with a spring-loaded bayonet mount is used to transfer sample blocks to and from the UHV chamber. Evacuation of the load-lock is accomplished with a Varian Turbo-V250 250 l/s turbomolecular pump (TMP) which provides pumpdown times to 10^{-7} Torr on the order of 30 minutes. This pump also provides preliminary roughing for the UHV chamber on those occasions that it is brought up to atmospheric pressure.

The UHV growth chamber consists of a stainless steel bell jar nominally 57 cm high and 35.5 cm in diameter. The centerline of the chamber is inclined 30° from the horizontal so that the sources all face upward toward the substrate. The UHV pumping system attaches to a vacuum flange which is located below the chamber. The load-lock attaches to a port on the side of the chamber. Opposite the load-lock is a liquid nitrogen (LN_2) filled cryoshroud which surrounds the substrate in the growth position. The sample

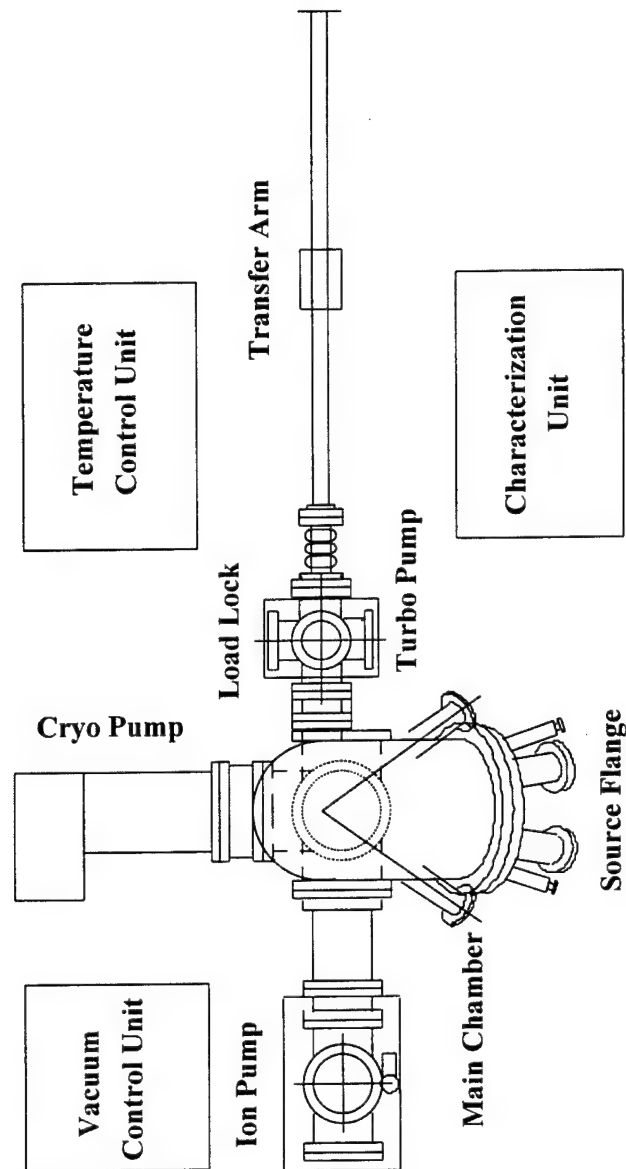


Figure 3.1 Schematic diagram of an MBE system. In addition to the main chamber, source flange, load-lock and pumping system, other important components include temperature control unit, vacuum control unit and characterization unit.

is held in the UHV chamber on an X-Y-Z manipulation stage to which is mounted the substrate heater and a bayonet mount for the sample block.

The UHV pumping system consists of a CTI-8 cryogenic (cryo) pump, a Varian 400 l/s ion pump and a titanium sublimation pump (TSP). The cryo-pump operates on the principle of Joule-Thompson expansion using He gas as a refrigerant. The expanding He gas cools a cold head to which is attached a baffle array and a high surface area absorber. The baffle array is cooled to 80 K and pumps by condensing species out of the vacuum space. The absorber is cooled to 15 K and pumps by either condensation or cryoabsorption depending on the species. The cryo-pump is a storage pump and as such loses pumping efficiency as the available pump area diminishes. To regenerate the pumping ability, the pump must be warmed and vented regularly. The ion pump ionizes incoming gas particles and then pumps by embedding the particle into a titanium (Ti) cathode or by reaction between the ion and Ti sputtered from the cathode by impinging particles. In the TSP, a thin layer of Ti is deposited on the walls of a LN₂ reservoir. This fresh layer of Ti is highly reactive with gas species such as hydrogen, nitrogen and oxygen and bonds them chemically to the reservoir surface thus removing them from the vacuum space.

The source ovens enter the chamber through a source flange at the bottom of the bell jar such that each oven has a nominal 29.3 cm source to substrate distance. Each oven has a molybdenum shutter which is used to block the flux from the oven until it is needed. The ovens are surrounded by a two layer cooling shroud which is kept at ~10° C by an H₂O: Ethylene Glycol (1:1) mixture cooled and recirculated by a Neslab (HX-200) water cooler. This shroud serves to isolate the ovens both to eliminate material cross contamination and thermal crosstalk. Oven temperatures can be controlled to better than 0.5° C using a Eurotherm microprocessor based controller. The nitrogen and hydrogen gas sources enter the chamber from side flanges located near the source flange. The

sources are mounted such that the hydrogen source is 34.8 cm from the substrate and the nitrogen source is 35.0 cm.

Chamber pressure is monitored using a Bayard-Alpert type ionization gauge mounted slightly behind the substrate position. The beam equivalent pressure (BEP) of the molecular flux impinging on the substrate can be measured by a similar ionization gauge mounted on a linear motion feedthrough. The linear motion allows the monitor to be positioned in front of the substrate for measurement and then retracted for growth. Residual gas composition in the chamber is monitored by a UTI 100C 1-400 a.m.u. quadrupole mass spectrometer.

3.1.2 Gallium Solid Source

Molecular beams of non-gaseous species are generated in standard MBE compatible Knudsen cells. These sources all contain a resistive heater assembly, a crucible to contain the source material and a thermocouple to monitor the source temperature (Figure 3.2). The thermocouple is monitored as part of a feedback loop with a Eurotherm 818 controller which allows temperature stability to better than 0.5°C. By precisely controlling the temperature of the source, one controls the vapor pressure of the source material in the cell and thus controls the material flux at the substrate. Figure 3.3 shows the output fluxes for this source for several operating temperatures.

The WVU MBE system uses an EPI-40-M modified filament cell to produce a flux of Ga atoms. This is a standard single filament source which provides additional heating in the area of the cell orifice. This so-called "hot-lip" is to prevent Ga droplet condensation at the edge of the crucible which could block the exiting flux or fall back into the molten Ga causing material spitting.

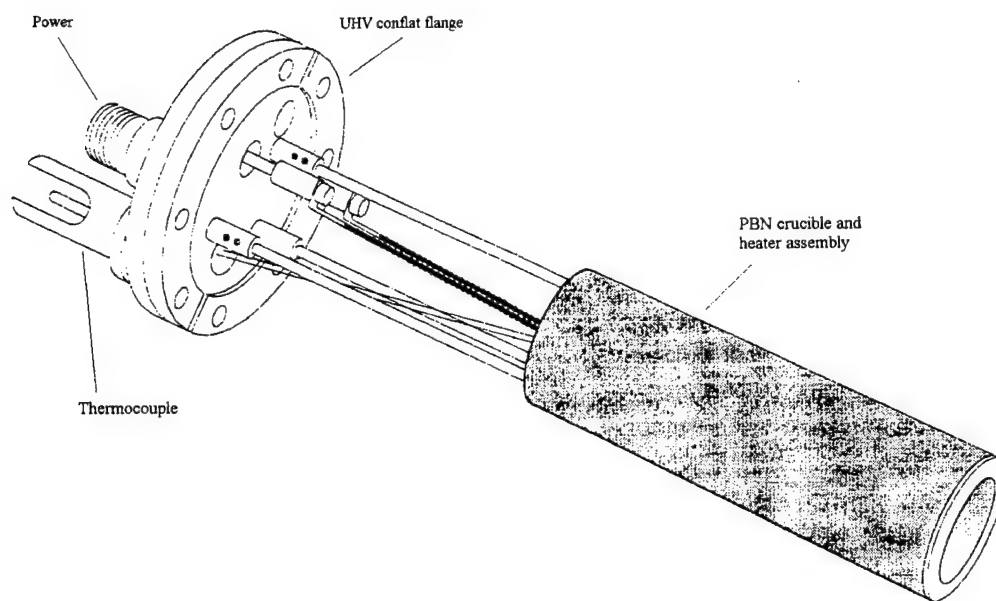


Figure 3.2 EPI-40-M Modified filament knudsen effusion cell.

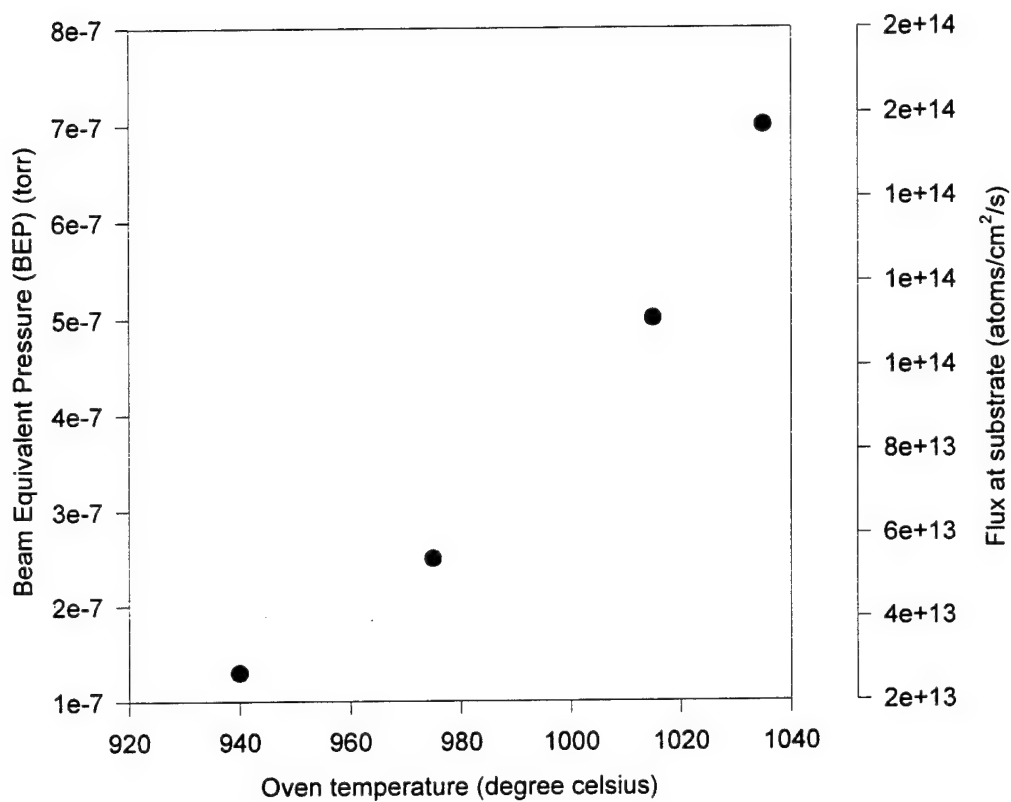


Figure 3.3 Beam equivalent pressure and equivalent atomic flux at the substrate for several Ga oven temperatures.

3.1.3 Atomic Hydrogen Source

The WVU MBE chamber is equipped with an EPI-AHS-L atomic hydrogen source. The EPI hydrogen source generates atomic hydrogen by thermally cracking molecular hydrogen on a heated tungsten filament. The filament is mounted axially down a tantalum tube inside the vacuum chamber. High purity hydrogen gas flows through a UHV leak-valve into the source and across the filament (Figure 3.4). The system is also equipped with an integral water cooling jacket to prevent excessive heating of the chamber. High purity deuterium gas can also be flowed into the source through the process gas manifold.

The source is operated by passing up to 10 amps through the filament in the presence of hydrogen gas. Typical operation in the course of this study applied 9.5 amps which corresponds to a power of 430 watts. This power produces a filament temperature of about 2200° C. (Figure 3.5 shows cracking efficiency vs. temperature) This corresponds to roughly 5% cracking efficiency with a 1×10^{-6} Torr beam equivalent pressure, and produces a flux of $8.6 \times 10^{14} \text{ s}^{-1} \text{ cm}^{-2}$ hydrogen atoms at the substrate.

3.1.4 Nitrogen Source

Due to the large (9.5eV) binding energy of the nitrogen molecule, low-temperature MBE growth of GaN cannot depend on the thermal dissociation of N_2 at the substrate. The chamber is therefore equipped with a radio frequency (RF) atom/radical beam source supplied by Oxford Applied Research, shown schematically in Figure 3.6, to provide active nitrogen. MBE growth using RF sources, like the one described here, and electron cyclotron resonance (ECR) sources is commonly referred to as plasma assisted (PA)MBE. Both types of sources have been successful in producing active nitrogen species with which GaN can be grown, however, ECR sources are known to generate a larger flux of nitrogen ions with energies high enough to damage the growing crystal and generate defects [10].

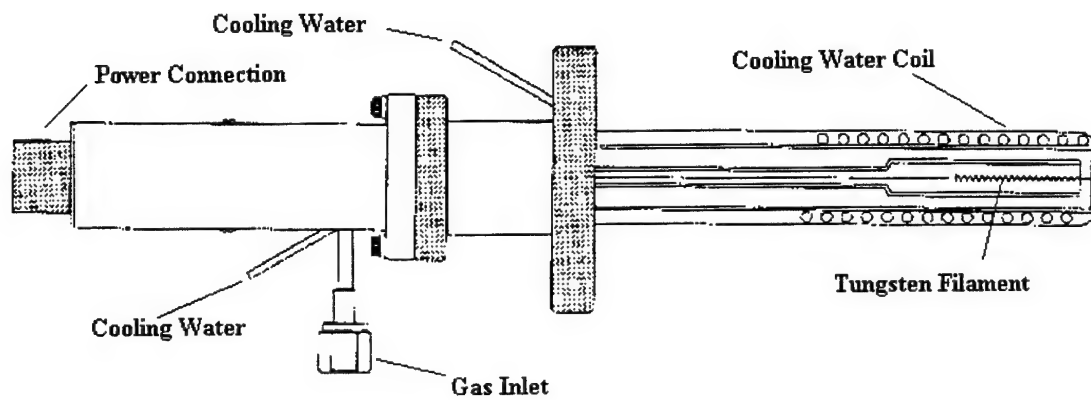


Figure 3.4 EPI atomic hydrogen source.

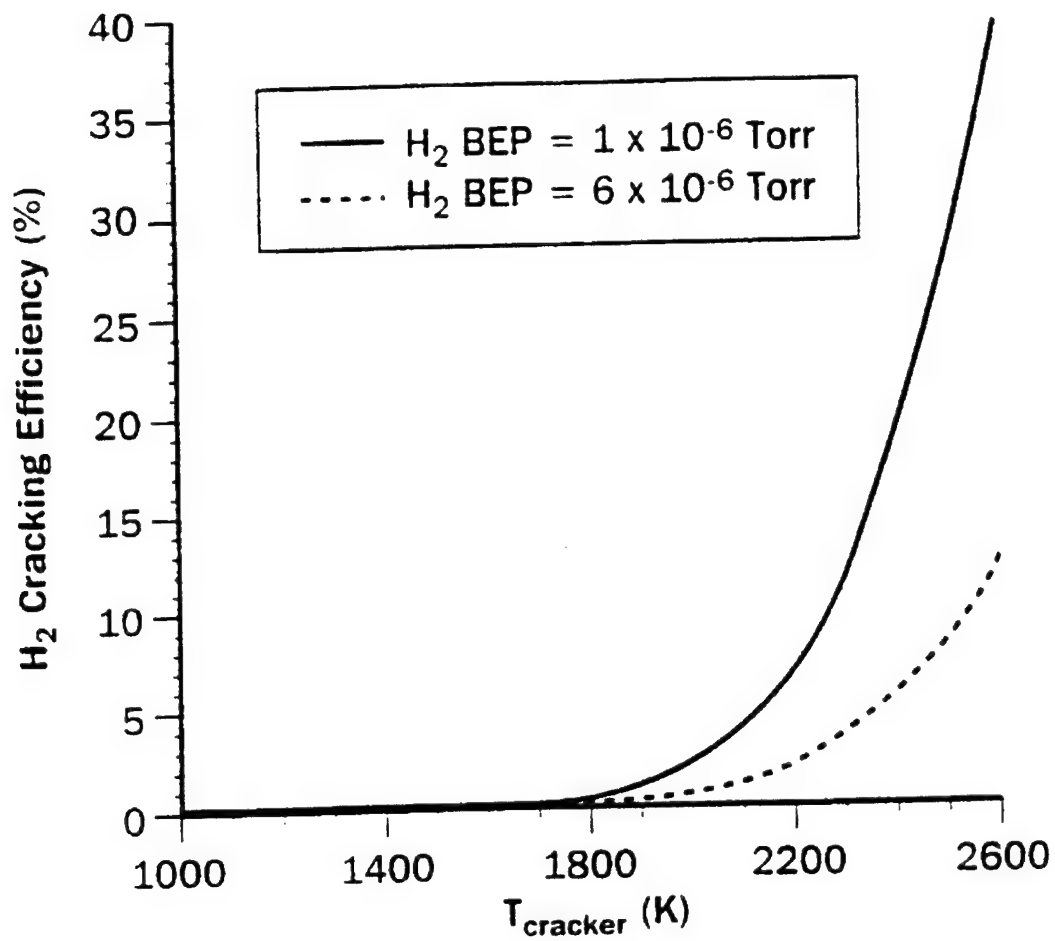


Figure 3.5 Cracking efficiency of atomic hydrogen source.

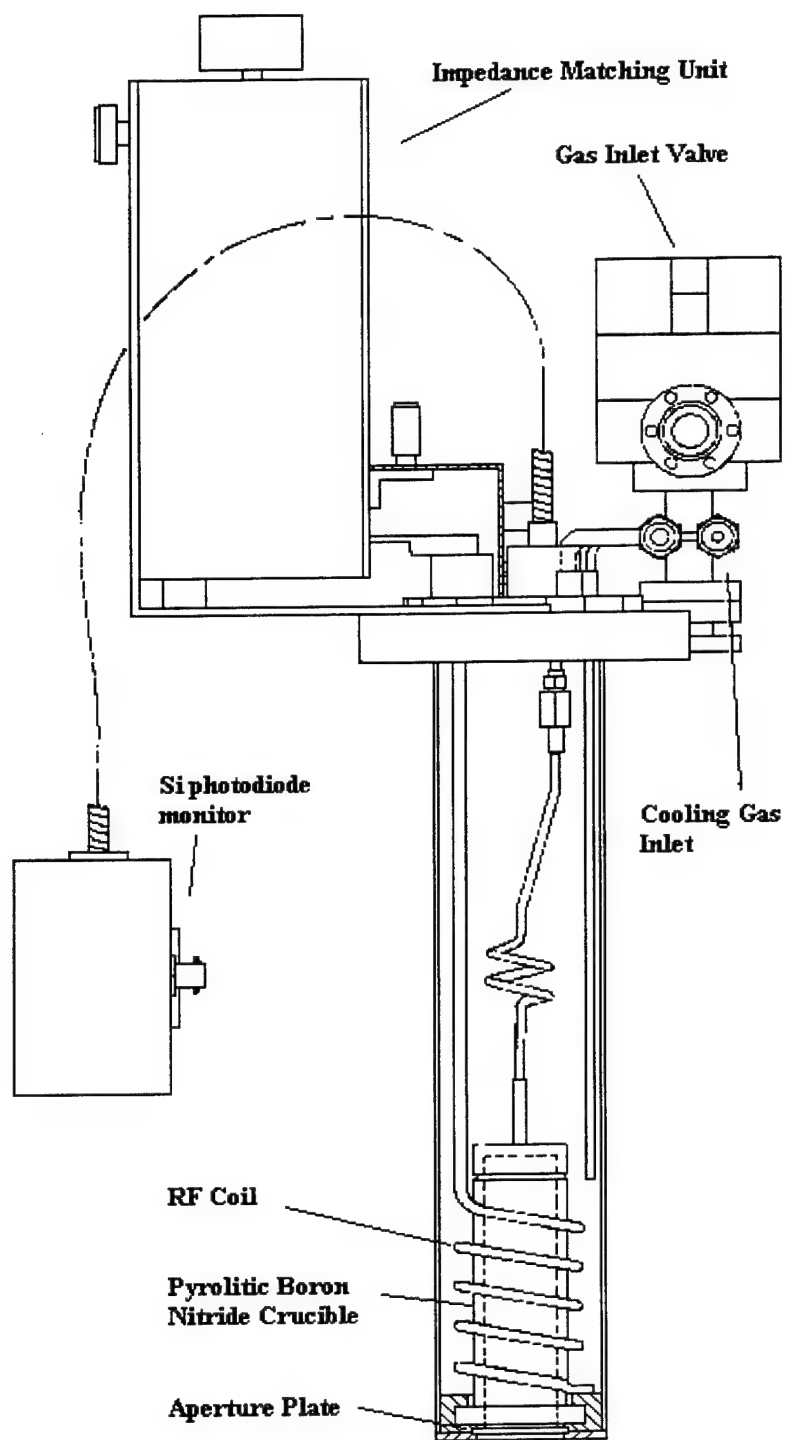


Figure 3.6 Oxford Applied Research CARS25 Radical/Beam source for the production of atomic nitrogen.

The Oxford CARS25 source is designed to fit into a standard MBE furnace port and consists of a pyrolytic boron nitride (PBN) crucible with a gas feed line on one end and a PBN aperture plate on the other. High purity (7N) nitrogen gas is flowed through the source during operation. An electrical discharge is established in the gas by an inductively-coupled 13.56 Mhz RF excitation. This discharge serves to generate a plasma in the N₂ within which, a fraction of the molecular nitrogen is dissociated. Under the continuous flow condition present in this source, a variety of species escape the source through the aperture plate into the vacuum environment. This resulting flux can conceivably contain a mixture of atomic and molecular nitrogen and ions of each. A thorough characterization of this flux is presented in chapter four.

The Oxford source can operate with up to 600W of RF power coupled into the plasma. The source is equipped with an optical port on axis with the plasma chamber through which it is possible to monitor the optical emission from the plasma with a photodiode and voltmeter. The source temperature is maintained by flowing dry, cryogenic N₂ gas, which is boiled off from a liquid N₂ dewar, around the plasma chamber and RF coil. The length of time for which a continuous GaN growth can occur is currently limited by the available cooling gas supply at a given RF power input.

PBN Aperture Plates for CARS25 Source				
Aperture	Application	Conductance (l/s)	# of holes	Diameter of holes (mm)
1	Doping	0.5	9	0.2
2	Doping	0.5	1	0.6
3	GaN Growth	10	37	0.5
4	GaN Growth	10	9	1.0
5	GaN Growth	10	1	3.0
6	GaN Growth	10	255	0.2

Table 3.1 Aperture plates available for the CARS25 Nitrogen source

The nitrogen source comes with two PBN aperture plates: one providing a low gas conductance (0.5 l/s) for doping fluxes, the other providing a higher conductance (10 l/s) for higher output flux. Additional plates with these conductances but varying geometries were manufactured by Angstrom Sciences (Pitt. PA) as part of the source characterization described later. Table 3.1 lists the attributes of the available aperture plates.

For a given aperture conductance and RF power, a stable plasma can be generated and maintained only within a narrow range of source inlet pressures.

3.2 Thin Film Characterization

3.2.1 Hall effect

Use of the Hall effect for semiconductor characterization provides a quick and inexpensive measure of a sample's electrical characteristics. Simple voltage measurements in the presence of a magnetic field provide important information on the density and sign of free carriers and also their mobility. Samples grown for this study were characterized in a room-temperature Hall effect system in the WVU MBE lab. A 6 kilo-gauss electromagnet is used to provide the DC field with a Keithley Model 220 programmable current source to provide the input current. A Keithley Model 7001 Switch box and Model 7065 Hall Effect Card provide the switching circuits used to average out contact and thermal effects. The Hall voltage is measured by a Keithley 197A digital voltmeter (DVM). System control and data acquisition are performed by an IBM PC over an IEEE-488 control bus (Figure 3.7).

Samples were measured using the Van Der Pauw 4-point geometry [20] which allows measurement of samples of arbitrary shape so that no post growth processing is required before measurement. Electrical contact was made by soldering wires directly to the GaN with indium. In general, all electrical contacts to semiconductors are rectifying

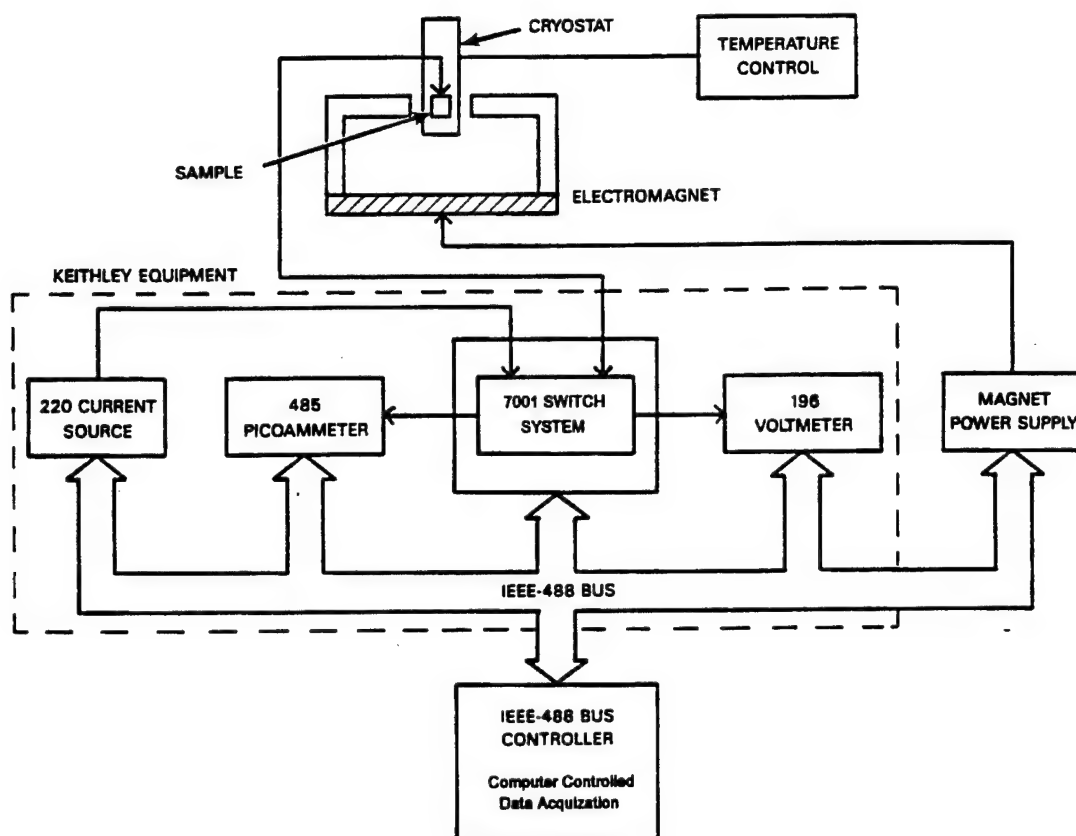


Figure 3.7 Schematic of a Hall measurement setup.

in nature so it is important to find an operating current which falls within the linear, ohmic region of the contact's current vs. voltage (I-V) curve. With this established, the computer is allowed to collect and average a large sample of voltages for each of the possible current and voltage contact permutations. These values are averaged and used to define the resistivity and Hall voltage. The Hall voltage is directly proportional to the Hall coefficient R_H . Once these values have been calculated, one can calculate the carrier concentration and mobility by the following equations.

$$n = (eR_H)^{-1}$$

$$\mu = \frac{R_H}{\rho}$$

3.2.2 Atomic Force Microscopy

Atomic force microscopy (AFM) was created in answer to the scanning tunneling microscope's inability to image a non conducting surface. AFM is a surface profiling technique where a sharp probe is scanned across the sample surface and the resultant probe deflection is measured [21,22]. This probe deflection can then be visualized either as the force between the sample and probe tip or as a change in the sample to probe tip distance which is converted into a relative height change. (See Figure 3.8) The AFM studies performed here were conducted in the lab of Prof. Naresh Dalal within the WVU Dept. of Chemistry by both the author and Dr. Michelle Richards-Babb using a Digital Instruments NANOSCOPE II AFM system. This open-air system was used primarily to assess large-scale surface morphology, at sub-optical resolution.

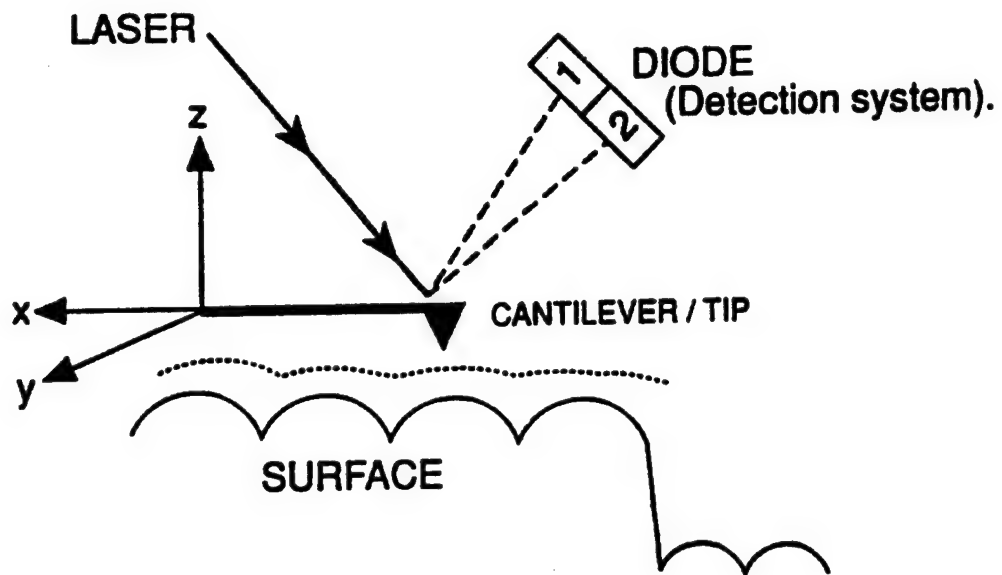


Figure 3.8 Schematic of the atomic force microscopy measurement system.

3.2.3 Photoluminescence Spectroscopy

Many types of semiconductor spectroscopies are based on the fact that a fraction of excess electrons and holes decay into lower level states by emitting a photon of light with the wavelength determined by the energy change between the initial and final states. By exciting a semiconductor with above-bandgap light to create excess holes and electrons and then analyzing the light emitted by the sample, it is therefore possible to identify radiative energy levels related both to the bandgap and to defects and impurities. PL studies were conducted by Prof. Nancy Giles in her laboratory within the WVU Physics Department on several samples grown during this work. Above bandgap excitation was provided by the 325nm line of a He-Cd laser focused to a nominal power density of 5 W/cm². PL was collected with a grating monochromator and a GaAs cathode photomultiplier. The input beam is optically chopped and the PL signal is run through a lock-in amplifier to increase the signal-to-noise ratio. For comparison to our results, a table of typical PL lines seen by other researchers is shown below [14].

Low Temperature GaN Luminescence Peaks	
Free excitons	3.474-3.475 eV
	3.480-3.481 eV
	3.49 eV
Free exciton LO replica	3.385 eV
Free exciton 2-LO replica	3.293 eV
Donor bound exciton (I_1)	3.44-3.47 eV
LO replica	3.377-3.378
2-LO replica	3.286 eV
TO replica	3.400 eV
Acceptor bound exciton	3.455 eV
Cd-acceptor bound exciton	3.454-3.455
LO replica	3.364 eV
2-LO replica	3.355 eV
Donor-acceptor	3.26-3.27 eV
LO replica	3.17-3.18 eV
2-LO replica	3.08-3.09 eV
3-LO replica	2.99-3.00

3.2.4 Transmission Electron Microscopy

A sample was characterized by transmission electron microscopy (TEM) by Dr. Linda Romano at XEROX Palo Alto Research Center (Palo Alto, CA). The sample was investigated by a combination of high resolution transmission electron microscopy, multiple dark field imaging and convergent electron beam diffraction techniques. The sample was cut parallel to the growth direction and then polished to a thickness of less than 5 μ m for electron transparency.

3.2.5 Secondary Ion Mass Spectrometry

Two samples were characterized by means of secondary ion mass spectrometry at Charles Evans and Associates (Redwood City, CA). By sputtering away sample material with an 8keV O_2^+ ion beam and monitoring the sputtered material with a Cameca magnetic sector mass spectrometer, a depth profile was developed for boron, potassium and lithium in the GaN samples.

3.2.6 UV Fluorescence Microscopy

This technique is similar to the photoluminescence spectroscopy described earlier in that UV light is used to excite lower energy radiative transitions in the sample. Instead of providing detailed spectral information, this technique provides spatial information for the source of the PL. Some spectral information is provided, however, by apparent color variations. Studies were performed using an Olympus BX60M microscope located in the WVU MBE lab. The microscope was equipped with a dichroic beam splitter to allow the output of a 100W Hg lamp to pass to the sample while allowing only visible and near-UV (380nm) light to be transmitted to the eyepiece or camera.

4. N₂ Source Characterization

For low temperature MBE growth of GaN, it is necessary to find a non-thermal method for breaking the 9.5 eV bond of molecular nitrogen. This has been attempted in a number of different ways. To date the most successful methods in the MBE arena have been to use either plasma sources, both RF and electron cyclotron resonance (ECR), or catalytic decomposition of ammonia on the GaN surface. The use of ammonia, though, brings in the problem of hydrogen, which can negate the benefits of the clean MBE environment over CVD processes. Plasma sources, on the other hand, allow control over the growth environment to be maintained. One question that remains, however, is the identification of the species produced by the source. It is likely that the flux contains both atomic and molecular nitrogen, possibly in a variety of excited states, and ions of each. Most researchers agree that it is neutral nitrogen which is responsible for GaN growth in MBE. Some groups have, however, discussed the possible benefits of low energy nitrogen ions [9] and also the detrimental effects of high energy ions [10]. It is important then that we understand what species are produced by our source in order to better define the growth parameters which lead to high quality film growth. In order to answer these questions, an optical and mass spectral analysis has been performed with the assistance of Prof. Charter Stinespring and his lab within the WVU Department of Chemical Engineering [23].

4.1 Characterization Technique

4.1.1 Mass Spectrometry

The Oxford Applied Research CARS25 source was installed in an UHV vacuum system in the WVU Chemical Engineering Department in direct line of sight with an Extranuclear Labs (Extrel) quadrupole mass spectrometer. The distance from the source

aperture plate to the ionizing volume of the mass spectrometer was approximately 8 cm. This is shown schematically in Figure 4.1. The chamber was pumped by one or more turbomolecular pumps (TMP) to a base pressure of roughly 10^{-9} Torr. The mass spectrometer itself was differentially pumped by a TMP to similar base pressures and maintained a pressure roughly two orders of magnitude lower than the test chamber during source operation.

The Extrel mass spectrometer consists of an ionizer, a set of electrostatic lenses, a quadrupole mass filter and an electron multiplier, shown schematically in Figure 4.2. A filament in the ionizer generates an electron beam which ionizes species within the ionizer volume. The ions created are then propelled into the lenses and mass filter by an electrostatic potential, henceforth called the ion energy, created between the ionizer and the mass filter. The ion energy potential also serves to repel external ions which have energies less than the set ion energy. Before entering the filter, the ions are focused into a beam by the lens array.

Once an ion beam has been created and focused, it needs to be separated and analyzed as a function of species mass. This is accomplished in the quadrupole mass filter. The Extrel mass filter consists of four conductive parallel hyperbolic cylindrical surfaces, arranged on the four corners of a cross, to which an RF voltage is applied. Opposing electrodes are connected together with one pair taking a potential

$$U(t) = U + U_1 \cos(\omega t)$$

and the other pair taking $-U(t)$. Ions entering the quadrupole with momenta roughly parallel to the z-axis undergo transverse oscillations in response to the ac and dc fields imposed in the x-y plane. For a given selection of U and U_1 , only ions with a specific mass to charge ratio will experience a stable oscillation and traverse the filter to be counted. Ions which do traverse the filter strike an electron multiplier generating a current which is measured as the signal intensity. By varying the ac and dc voltages in the filter

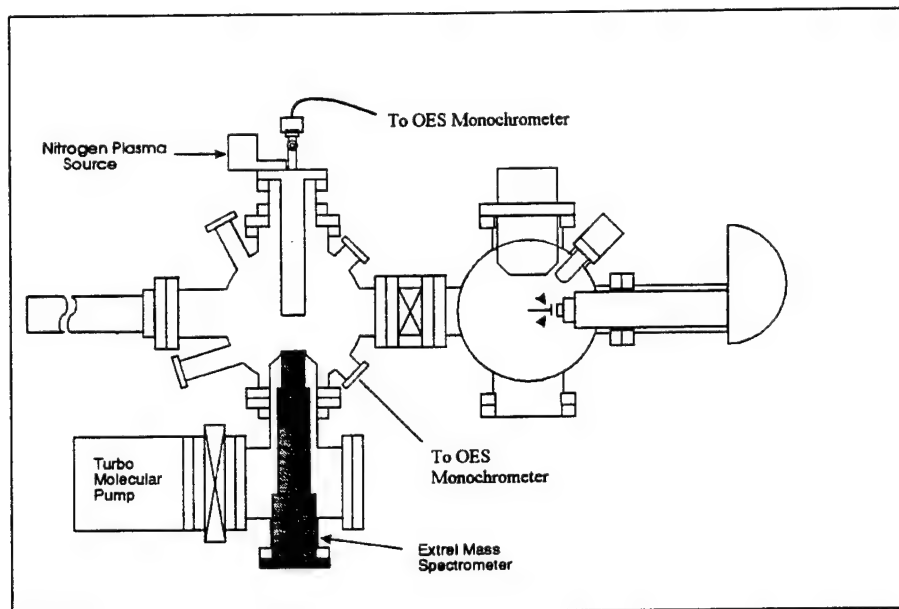


Figure 4.1 Schematic of nitrogen source characterization chamber. Source is mounted down the mass spectrometer line-of-sight. The distance from the source to the mass spec. is roughly 8 cm.

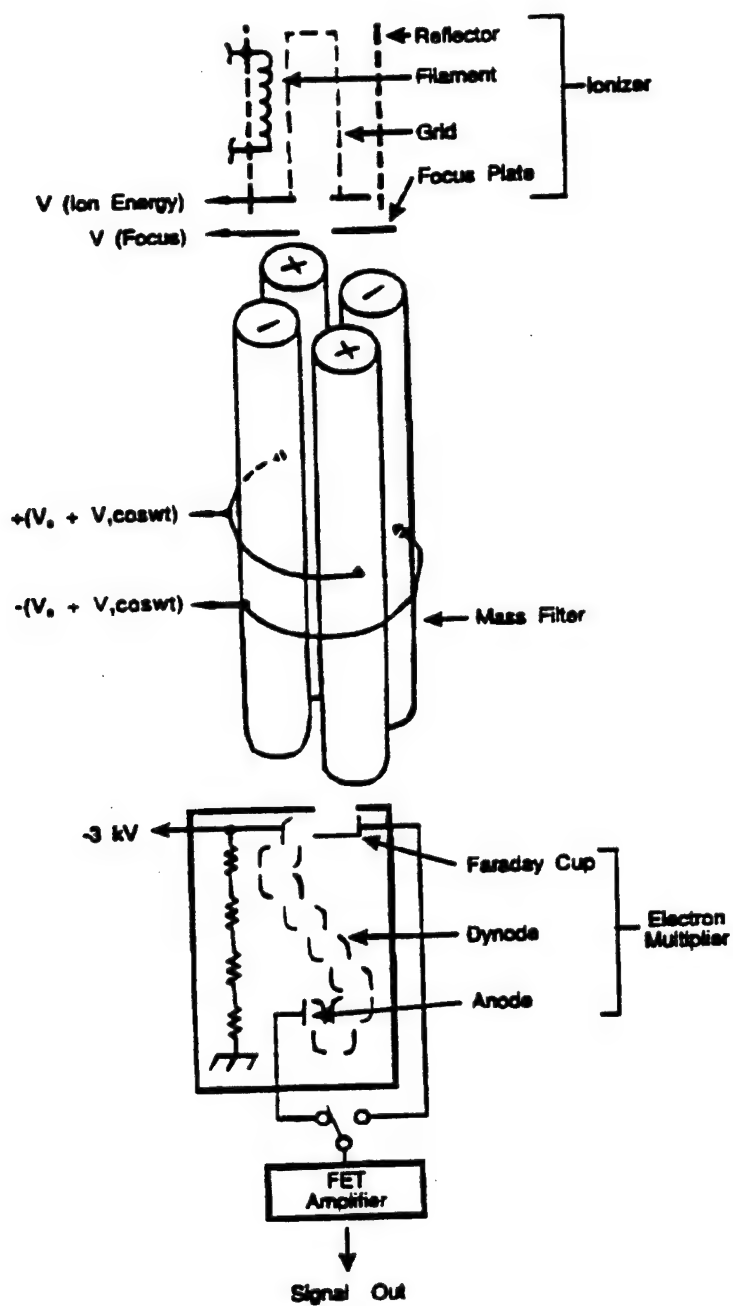


Figure 4.2 Schematic diagram of QMS system.

and monitoring the signal intensity, it is possible to generate a spectrum for a range of mass to charge ratios.

The fact that the quadrupole filter measures the charge to mass ratio and not the mass directly immediately poses one problem for the current research. The purpose of this characterization was to differentiate between N and N₂ and ions of each in the source output flux. For N⁺ and N₂⁺, this is easily accomplished but, if N₂⁺⁺ should be present, it would be indistinguishable from N⁺. For ions created in the mass spectrometer (neutrals in the source output flux), it was sufficient to keep the energy of the ionizer electron beam no higher than 20 eV. This energy is sufficient to efficiently ionize all species of interest without being high enough to dissociate or doubly-ionize the N₂.

Figure 4.3 shows the transmission function for the quadrupole filter as a function of ion energy for both atomic and molecular nitrogen. This figure was produced, with the source turned off and a fixed background of N₂, by adjusting the electron energy of the ionizer up to 70 eV. The resulting electron impacts have enough energy to dissociate some of the molecular nitrogen. A mass spectrum was then compiled for both the atomic and molecular peaks as a function of ion energy. The atomic nitrogen transmission peak is shifted to slightly lower energy compared to the N₂. We believe that this is due to N⁺ ions acquiring a higher velocity than the N₂⁺ in the ionization process. For subsequent analysis, the transmission functions for N and N₂ are taken to be the same. Similar transmission functions were obtained for each source aperture and operating pressure and were later used to correct the energy distribution data.

Since the source is likely generating an ionic component to its flux, it was necessary to be able to discriminate the ionic contribution from the source and that generated in the mass spectrometer. This was possible by operating the mass spectrometer in two modes, low emission and high emission. In low emission operation, the filament current is reduced to 0.018 mA effectively stopping ion production in the mass spectrometer ionizer. Any signal measured in the system must then be due to ions

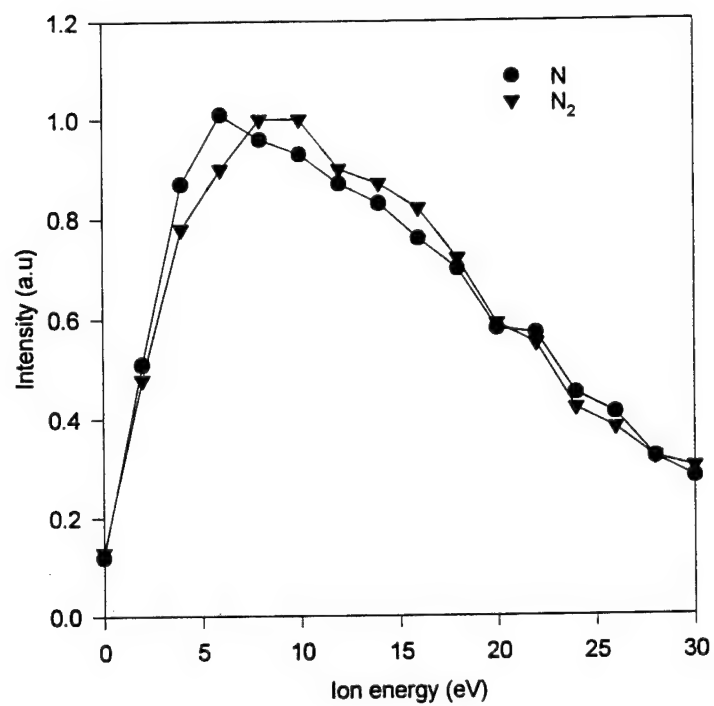


Figure 4.3 Mass filter transmission function for N and N₂

originating outside the ionizer. In high emission operation, the filament current is turned up to 1 mA so that ions are generated in the ionizer. The measured signal is now a combination of the signals from external and internal ion sources. By subtracting the low emission data from the high, one gets the signal due to neutrals in the source flux. This data is then adjusted to the transmission function and scaled to the ionization fraction of the particular species.

4.1.2 Optical Emission Spectroscopy

The CARS25 source is equipped with an optical emission monitor which uses a fiber optic cable to sight down the axis of the source to monitor plasma light output. It has been suggested that this monitor might be used as a process indicator to fine tune the output for maximum atomic nitrogen flux. Source characterizations performed by other labs [24-28] have used spectral analysis of this output to determine the composition of the output flux. It was hoped that the output of the optical monitor could be matched to the mass spectral data to provide some useful control information.

The principle behind optical emission spectroscopy is that an excited molecule will eventually radiatively de-excite emitting a photon with a wavelength dictated by the change in energy between the excited and 'ground' states. Typically, these energy states are unique and provide a fingerprint for the identity of the radiating species. By monitoring the source's optical output with a monochrometer, it is possible to extract these 'fingerprints' and see how they vary as a function of the source operating conditions.

Light output was monitored via a fiber optic cable initially connected to the optical port at the rear of the plasma source. This was later moved such that the fiber viewed the exit aperture at the front of the source (Refer back to Figure 4.1). In normal operation, this light is monitored by a silicon photodiode and the integrated intensity is displayed on a digital voltmeter (DVM). For this study, the light was passed into an Instruments SA, Inc. HR-320 Spectrograph. The HR-320 has a 0.32m focal length and

was operated with a 1200 grooves/mm grating. A stepper motor under computer control drives the spectrograph to scan through a wavelength region. The spectrograph output was monitored by a Si photo-voltaic detector, which was read by computer through an IEEE-488 interface on a Keithley 197 DVM. Scans could then be saved for later analysis. A typical scan is shown in Figure 4.4.

Vaudo *et al.* [26,27], performed high resolution spectral scans on a similar nitrogen source and showed conclusively that the peaks at 745, 821, and 869 nm are due to transitions in atomic nitrogen. These peak assignments were used here for the spectral analysis. Peaks were fit using Jandel Scientific Peakfit non-linear curve-fitting software. All peak heights were normalized to the 773 nm molecular line so that a comparison could be made between the relative molecular and atomic peak intensities.

4.2 Results

Analysis was performed on four source aperture plates described in section 3.1.4. Apertures 1 and 2 are typically used for doping applications requiring a low flux. Apertures 3 and 4 are representative of the aperture used for GaN growth.

During typical operation of this source, a silicon photodiode monitors the integrated light intensity from the plasma chamber. The output of this detector is displayed on a digital voltmeter (DVM) and used as a primary indicator of plasma condition. Figure 4.5 shows representative plots of this photodiode response for the high and low conductance apertures used on this source. For a given RF power and aperture plate, a stable plasma can be maintained only within a narrow window of pressures. In general, the width of this existence window increases with applied power. At a fixed power, for source pressures on the low end of this existence window, the photodiode response increases linearly with increasing source pressure. The response eventually flattens out becoming relatively constant. At high enough source pressures, the response falls off again, decaying until the plasma becomes unstable.

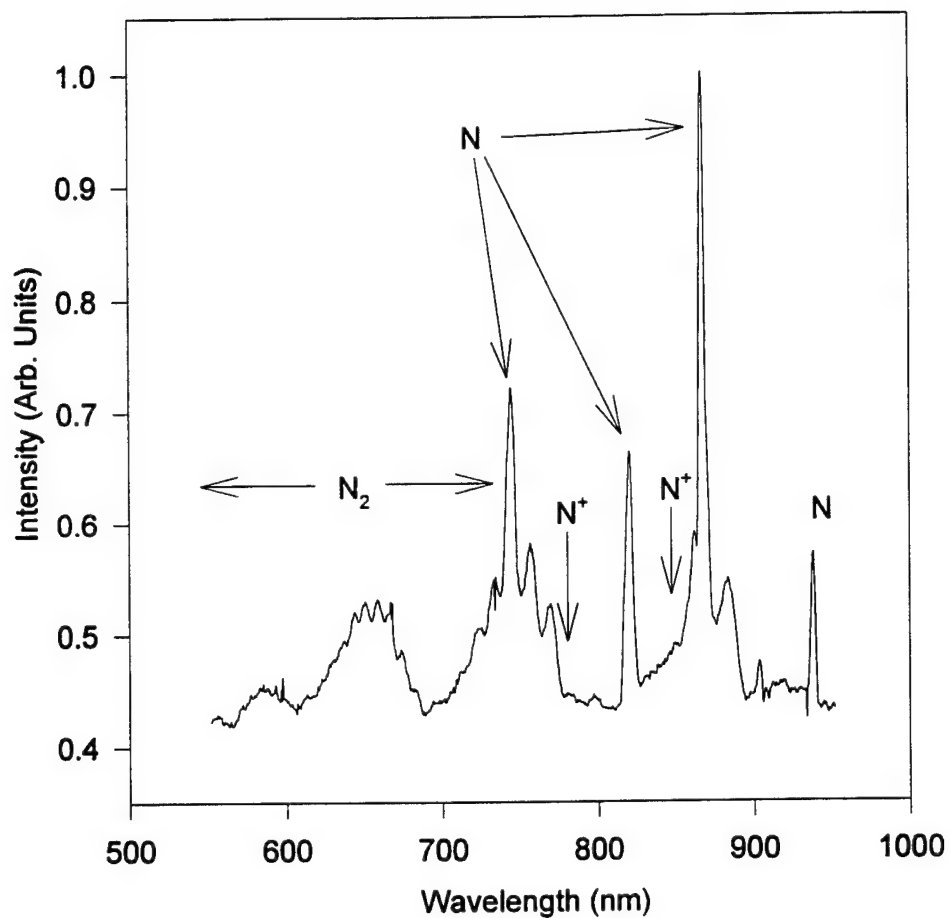


Figure 4.4 Typical spectrum of the optical emission from the CARS25 nitrogen source during operation. The labeled peaks have been related to transitions in atomic nitrogen.

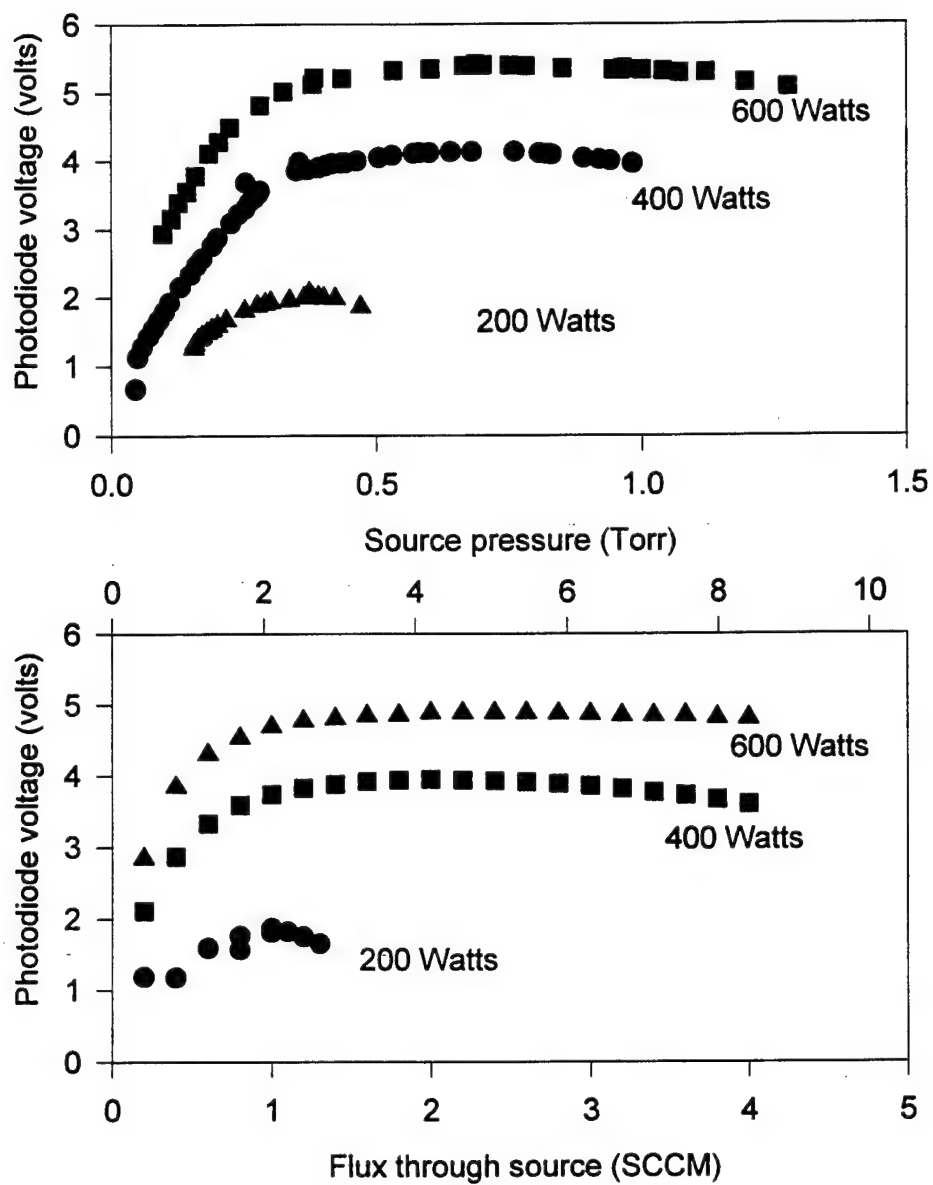


Figure 4.5 Photodiode response as a function of source pressure (Nitrogen flow for the lower plot) The upper plot is characteristic of the doping aperture, while the bottom is characteristic of the growth aperture.

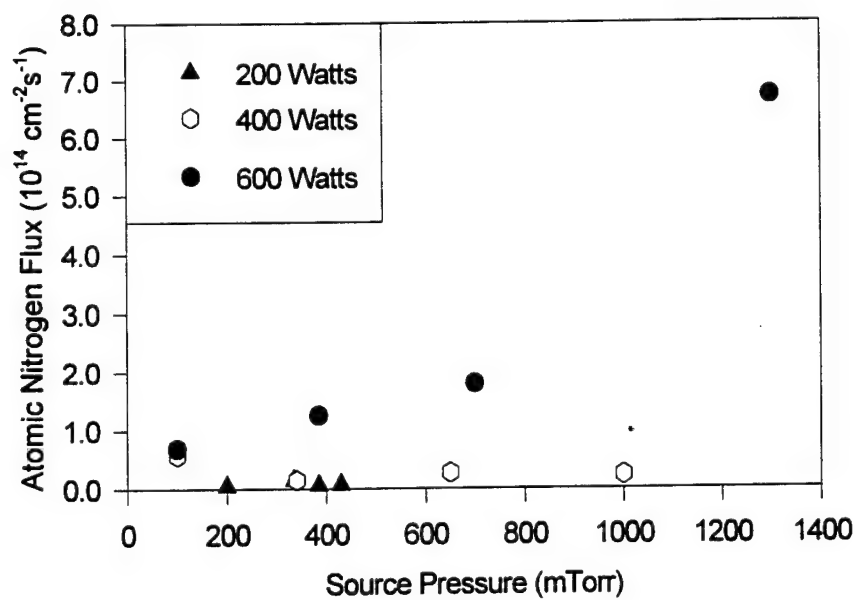
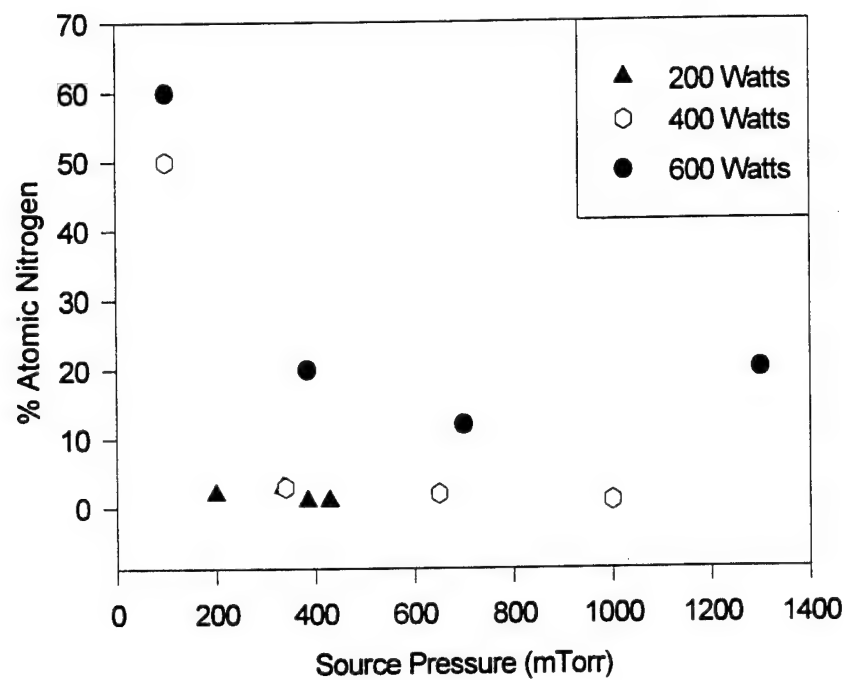


Figure 4.6 Summary plot of nitrogen source mass spectra for doping aperture #1(top). Bottom plot shows atomic nitrogen flux at substrate. Relative error in these values is estimated to be approximately 20%.

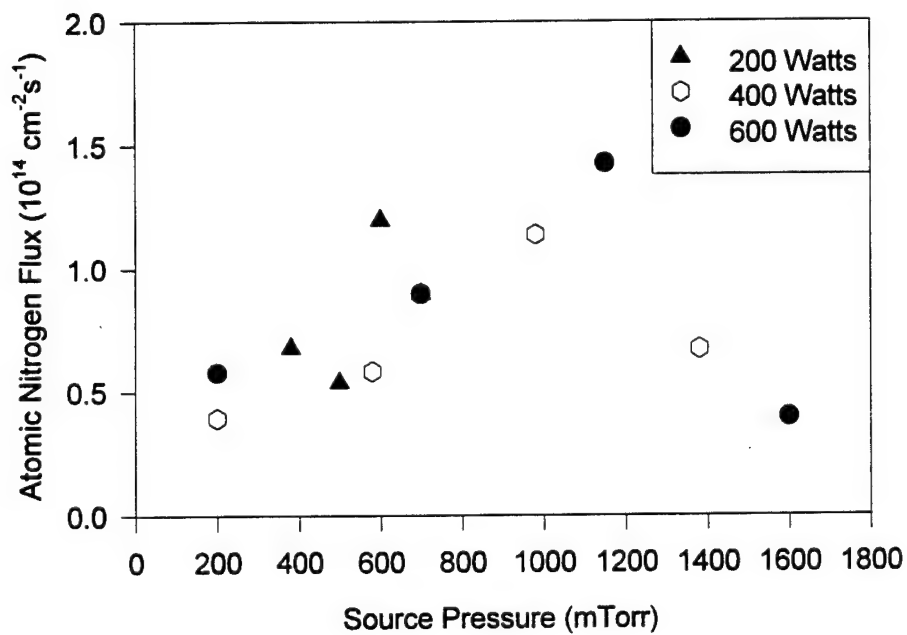
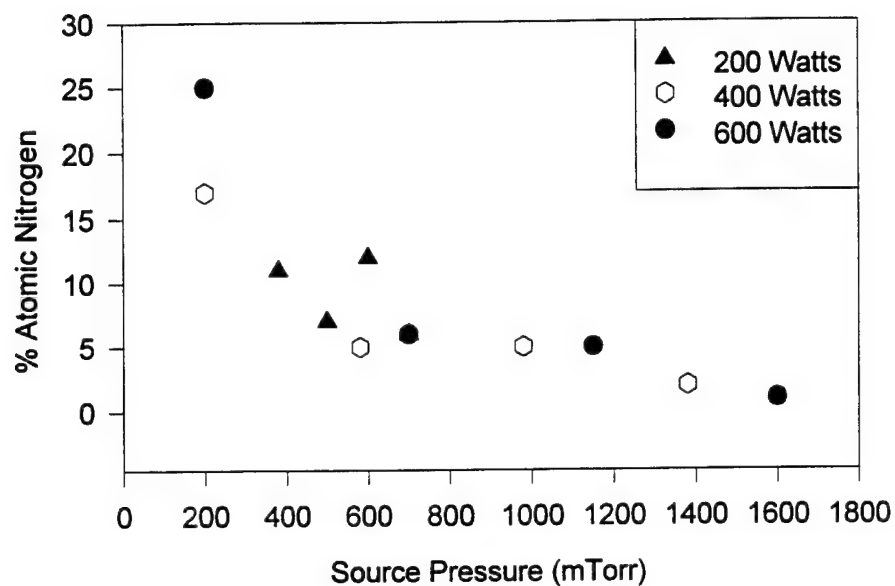


Figure 4.7 Summary plot of nitrogen source mass spectra for doping aperture #2(top). Bottom plot shows atomic nitrogen flux at substrate. Relative error in these values is estimated to be approximately 20%.

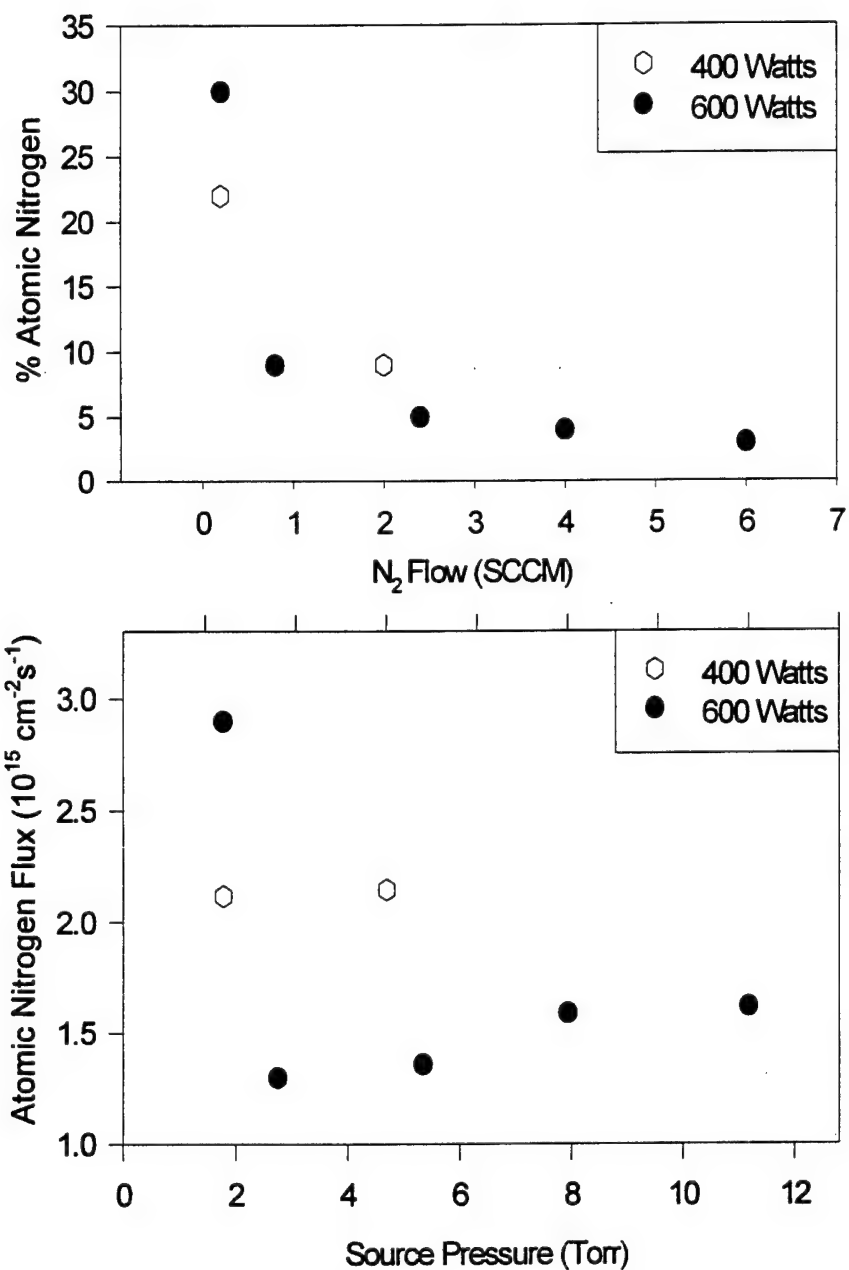


Figure 4.8 Summary plot of nitrogen source mass spectra for growth aperture #3(top). Bottom plot shows atomic nitrogen flux at substrate. Relative error in these values is estimated to be approximately 20%.

Figures 4.6-4.8 show the results of the mass spectral analysis for apertures 1-3. In all cases, the percent conversion to atomic nitrogen decreases with increasing source pressure, at constant power. However, when scaled to actual output flux, the atomic nitrogen flux increases with pressure. Increasing power, on the other hand, is always seen to increase the fraction converted over the range investigated. From this a consistent picture begins to develop where increasing the applied power increases the energy available for molecular dissociation. Increasing the source pressure, however, increases the density of molecules and thus the probability that two atoms will collide and recombine in the source.

All of these analyses relied on the source output flux containing ions with energies lower than 50 eV. External ions with energies higher than this could not be rejected by the ionizer system and made extraction of the neutral flux difficult, if not impossible. Several points had to be rejected from these figures because a high energy ion component caused a severe overestimation of the neutral atom content. Data for aperture #4 had to be rejected entirely because of a large flux of high energy ions coming from the source. It was impossible to extract useful information for this aperture.

From this result, it appears as though ion recombination is enhanced in the boundary layer region around the aperture plate exit holes. Larger exit holes, as designed for aperture #4, allow more ions to go through without traversing the boundary layers and being neutralized. In order to minimize ion content in the output flux, smaller holes are necessary. For subsequent studies, an aperture plate has been made with the conductance of aperture 3 but with 0.2 mm holes, in the hope that this will reduce the ion flux.

Figures 4.9 and 4.10 show data from the peak fits performed on the OES data for aperture 1. All data in these figures have been normalized to the 773 nm molecular line intensity. In Figure 4.9, the OES output for the atomic nitrogen lines tracks with the mass

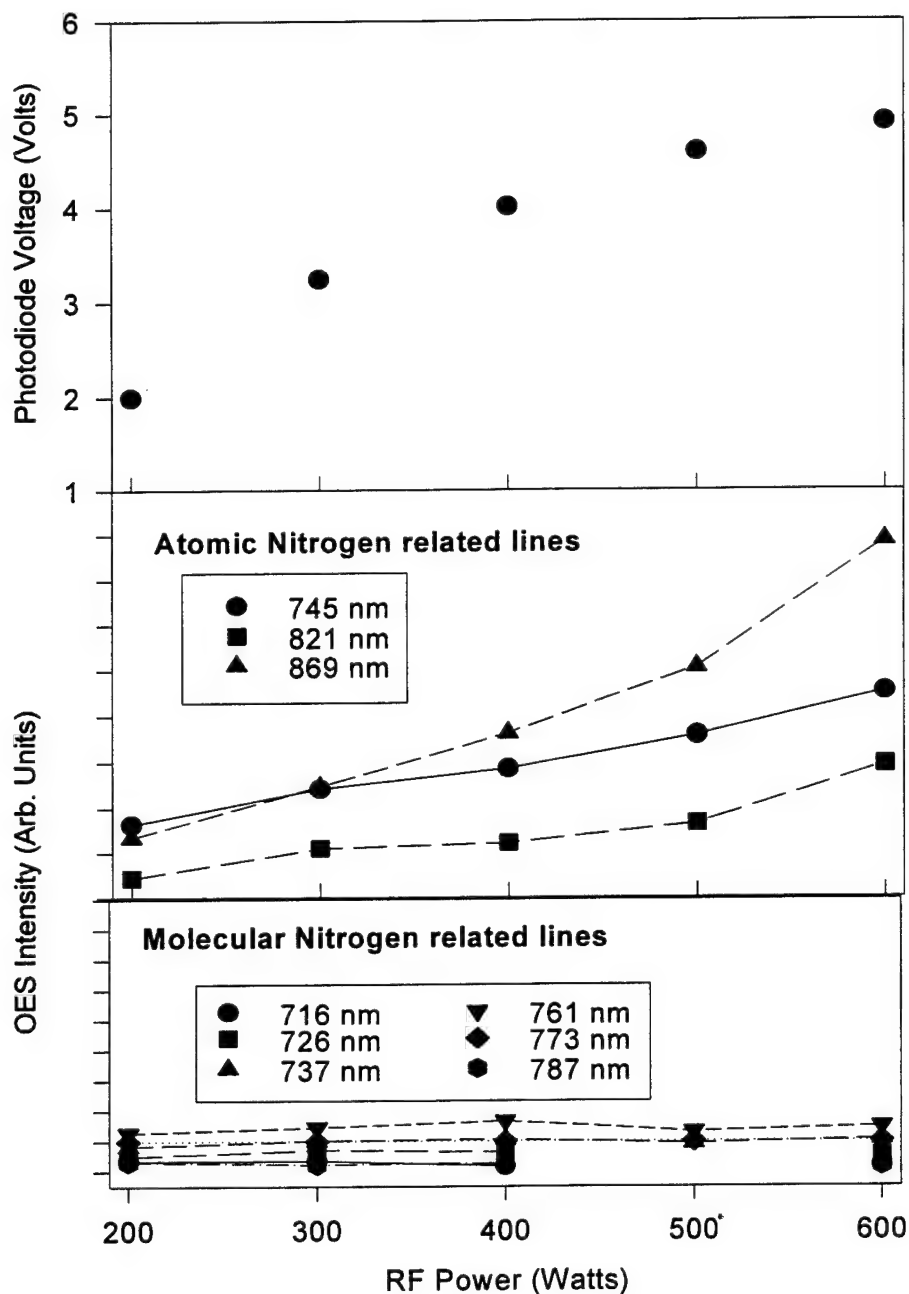


Figure 4.9 Representative plots of atomic(middle) and molecular(bottom) line intensities , normalized to the 773 nm line intensity, from OES peakfits at fixed source pressure. Top graph is the integrated intensity from the photodiode system included with the source. Similar results were seen for all apertures studied.

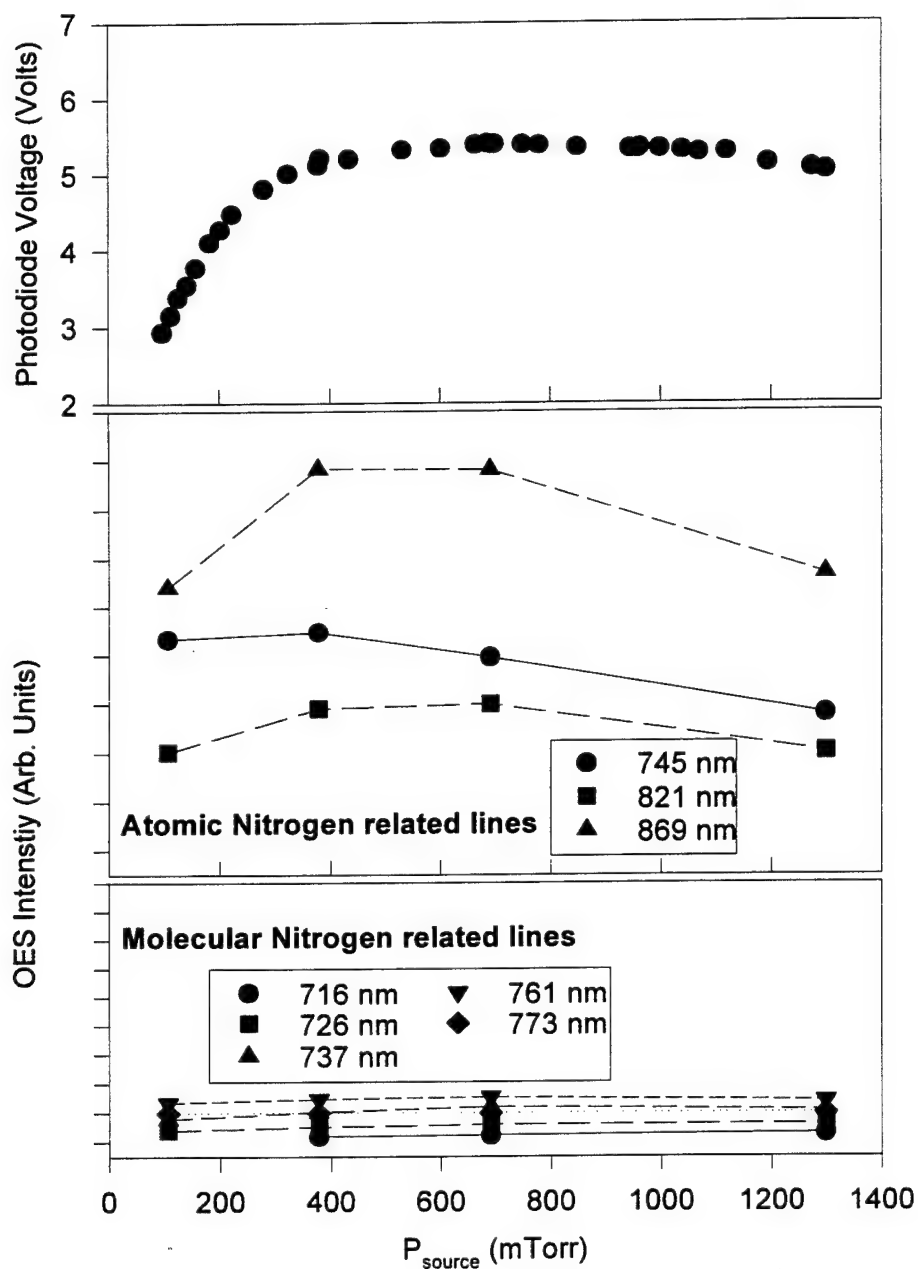


Figure 4.10 Representative plots of atomic(middle) and molecular(bottom) line intensities, normalized to the 773 nm line intensity, from OES peakfits at fixed source power. Top graph is the integrated intensity from the photodiode system included with the source. Similar results were seen for all apertures studied.

spectroscopic data: as the RF power increases, so do the atomic peak intensities relative to the molecular peaks. Note that the actual intensity of both line-sets increased with power. In Figure 4.10, the atomic lines are seen to increase and then fall off again with increasing pressure. In both cases, the atomic lines seem to track well with the integrated intensity displayed at the top of each figure. These same optical trends were seen for all of the apertures tested. In all of these cases, it appears that the optical emission from the atomic lines tracks reasonably well with the integrated emission from the plasma, however, its relationship to the actual output flux content is difficult to ascertain.

For all but aperture 4, the ionic contribution to the source flux was low, about 0.03%. Note, however, that this corresponds to 2×10^{13} ions/cm²/sec and could generate a volume defect level of 10^{17} cm⁻³ for typical growth rates of 0.1 μ m/hr. Maximum ion energies were mostly below 25 eV. Ion energies for aperture 4 were greater than 50 eV, which is generally considered to be entirely detrimental to GaN growth.

5. Nucleation and Growth of GaN

As discussed in section 2.2, growth of GaN on sapphire typically occurs in a three-dimensional mode, beginning with the nucleation of isolated islands that maintain their individual character during subsequent growth. Such columnar growth, due primarily to the large lattice and thermal mismatch, leads to a rough epilayer containing a high concentration of defects, which are mainly threading dislocations. The use of a low temperature GaN or AlN buffer layer, grown at 400-650 °C and annealed at a higher temperature before resuming growth, has been shown to dramatically improve layer morphology and electrical characteristics in the grown layer.

5.1 Nucleation and Buffer Layers

The initial nucleation layer can dictate much of the subsequent growth quality in heteroepitaxial growth. To investigate the nucleation and initial growth of GaN on sapphire, a series of layers was grown under varying conditions with film thicknesses ranging from 100Å to 0.4µm.

All films were grown on 1 cm² C-plane (0001)sapphire substrates purchased from Union Carbide Crystal Products. Substrates were degreased in heated baths of trichlorethylene, acetone and methanol and then etched for ten minutes in a phosphoric/sulfuric (1:3) acid mixture heated to 160 °C to remove residual polishing damage. Following this etch, the substrates were rinsed in de-ionized water (characterized by a nominal 18MΩ resistance) and blown dry with high-purity N₂ gas. Substrates were mounted to a molybdenum block with graphite (Aquadag) and a molybdenum mask, and placed into the chamber load-lock. After pumping out the load-lock, the sample block was transferred to the growth chamber and heated to 100 °C for one hour to outgas the sample and the Aquadag. Substrates were then heated to 730 °C

and exposed to an atomic hydrogen flux, from the thermal cracker described earlier, for twenty minutes to remove surface carbon and oxygen contamination.

Early in this study, several layers were grown using nucleation conditions reported by others [29,16]. Following the prevailing wisdom of GaN/sapphire growth, substrates were "nitrided" by exposure to a nitrogen plasma prior to growth. This supposedly produces a thin layer of AlN that then assists in providing strain relief for the subsequent film growth. Growth of a buffer layer on this surface resulted in a fine-grained ($<100\text{\AA}$) island size seen under AFM (Figure 5.1). Growth of an initial layer at low temperature ($450\text{ }^{\circ}\text{C}$), without nitridization, followed by an anneal at $660\text{ }^{\circ}\text{C}$ also resulted in a similar small grain size (Figure 5.2). With grain diameters of this size, the resulting films are virtually guaranteed to have dislocation densities on the order of 10^{12} cm^{-2} . This level of dislocation density has been observed in MBE material grown in many laboratories [9]. If the nucleation domains could be made larger, the effect would be to reduce this intrinsic defect concentration. As such, a series of growths was conducted to optimize the growth parameters for the nucleation layer to increase the nucleation domain (island) size. Domain size distributions were obtained by collecting ten AFM micrographs at points distributed around the sample. The islands were then approximated as circles by visual comparison with a template. From this, a histogram of occurrence frequency vs. diameter was obtained. The distributions were adequately represented by a poisson distribution, and the mean value was found by a least squares fit to the distribution (Figure 5.3). The fact that the island diameter followed a poisson distribution indicated that the nucleation process was statistical in nature.

Figure 5.4 is a compilation of these measurements for various growth conditions. As the substrate temperature is increased for a given Ga and nitrogen flux, the nucleation island diameter is seen to increase somewhat linearly up to some maximum island size after which island diameter decreases with increasing temperature. Increasing the Ga flux increases both the maximum island size and the maximum temperatures at which island

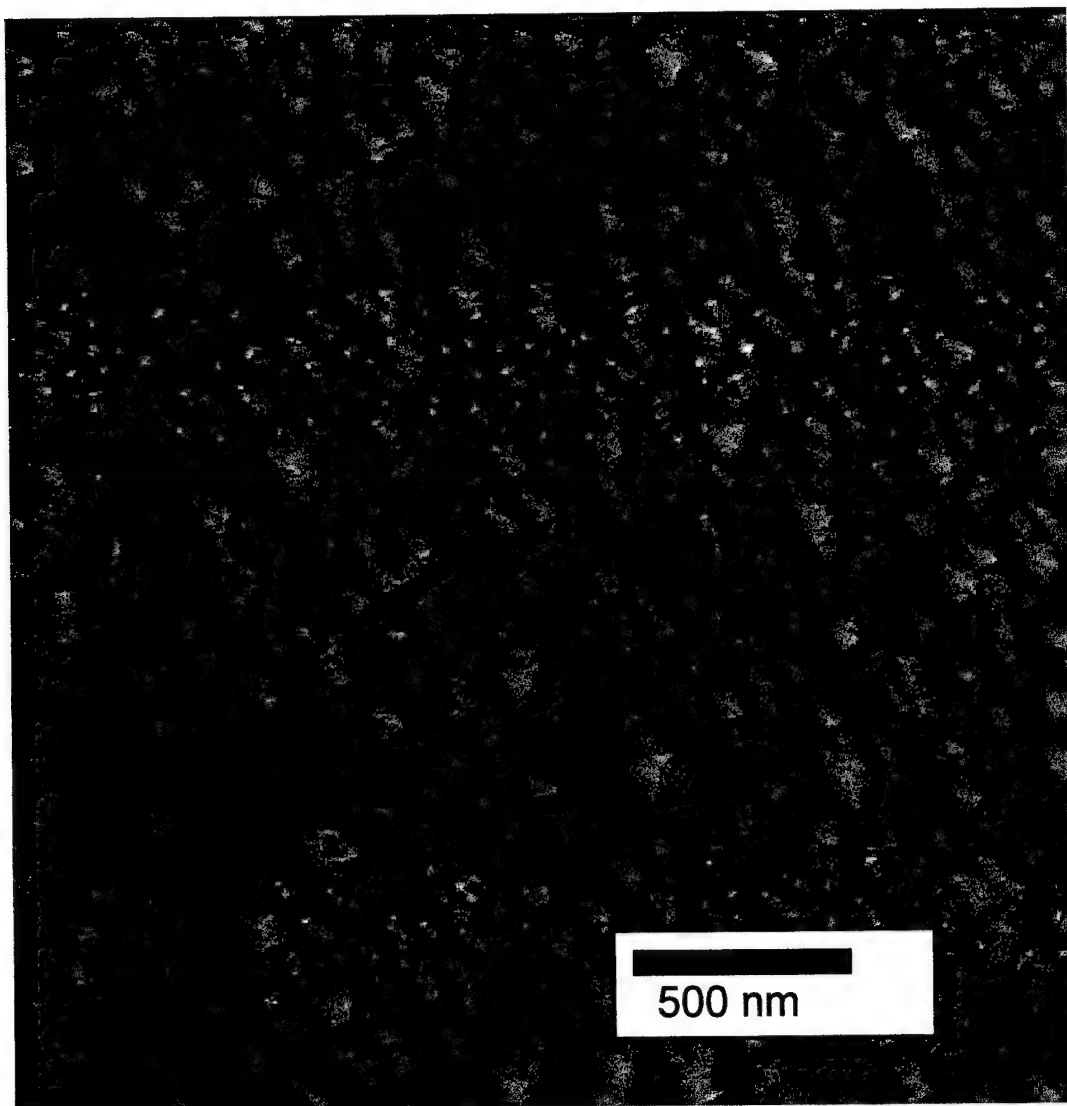


Figure 5.1 AFM micrograph of sample 9555 grown at 630C substrate temperature, 2.5×10^{-7} tor Ga BEP, 4 SCCM N_2 and 500 watts. Sapphire was exposed to nitrogen plasma prior to growth.

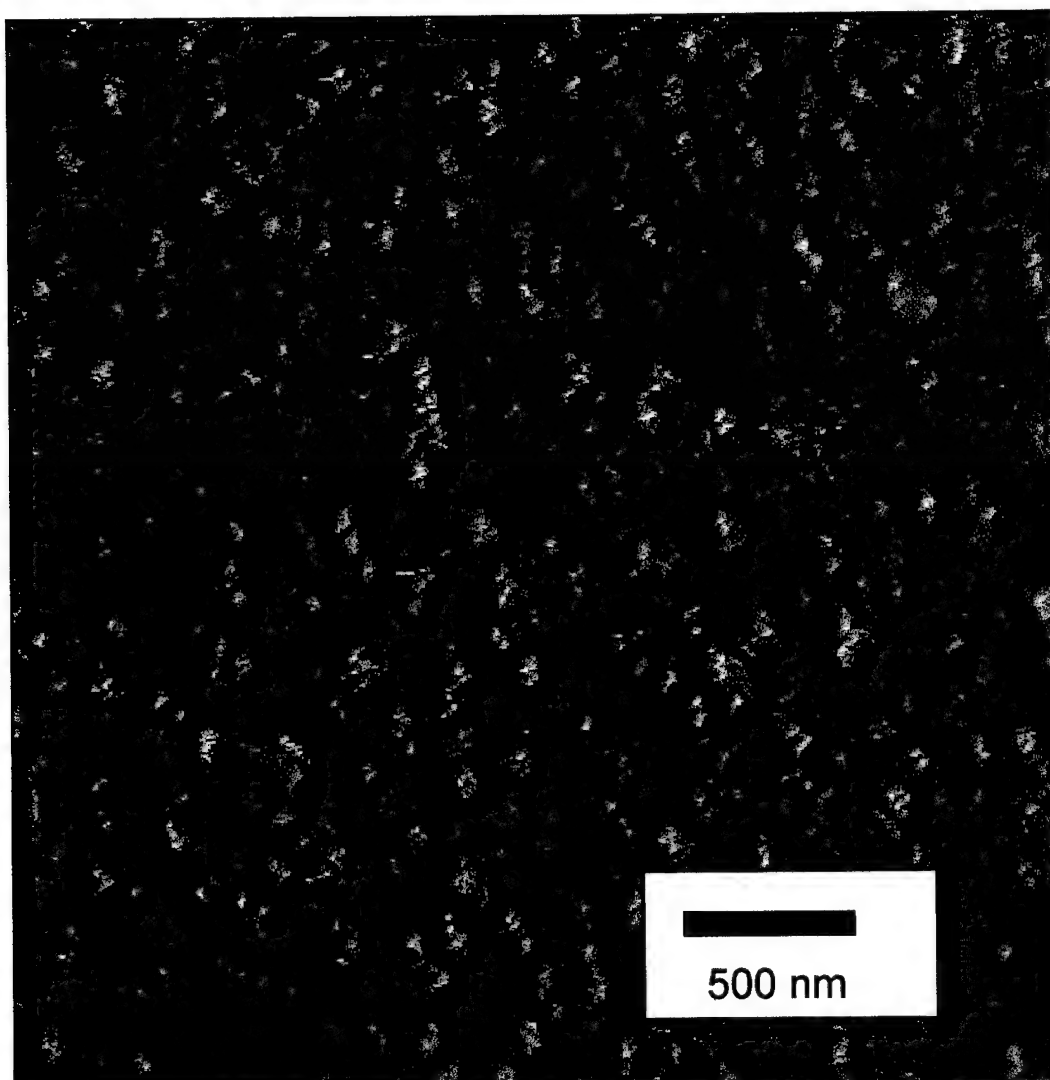


Figure 5.2 AFM micrograph of sample 9558 grown on a 50 angstrom buffer layer grown at 450C and annealed at 630C. Epilayer grown at 630C substrate temperature, 2.5×10^{-7} torr Ga BEP, 4 SCCM N_2 and 500W.

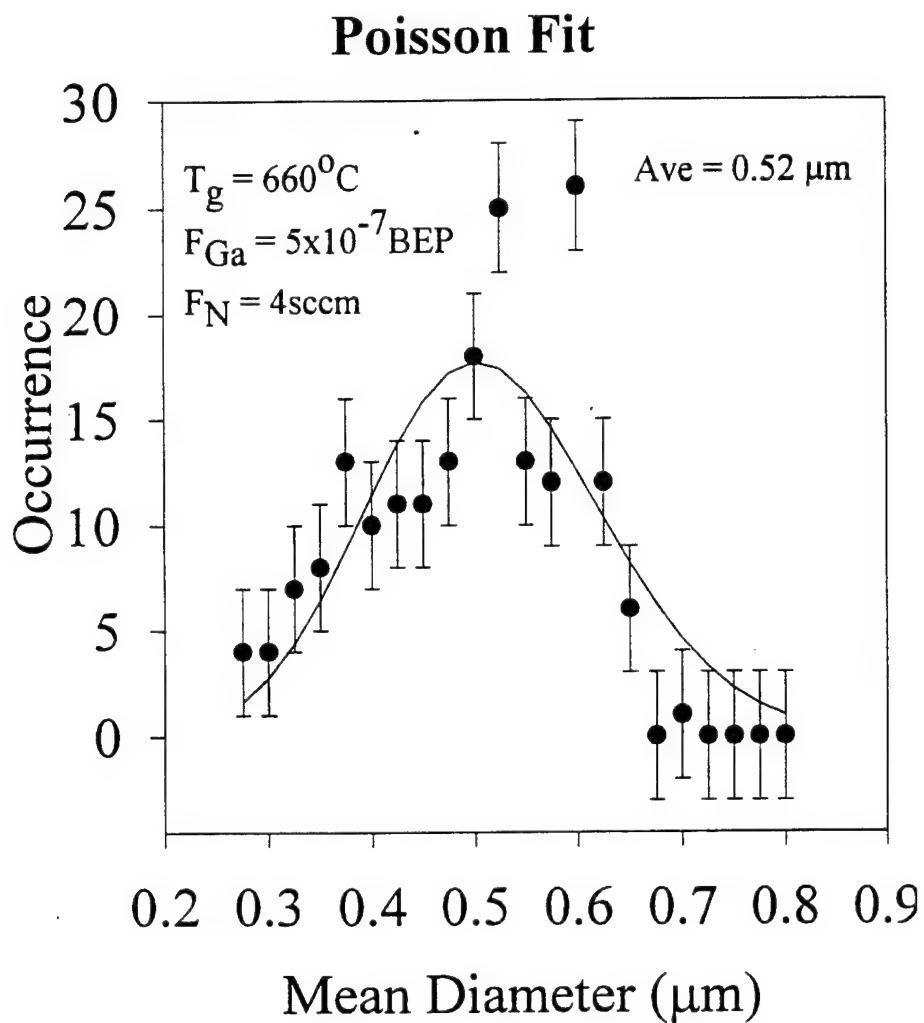


Figure 5.3 Representative fit of island diameter distribution to poisson distribution. Mean island sizes were extracted from similar fits.

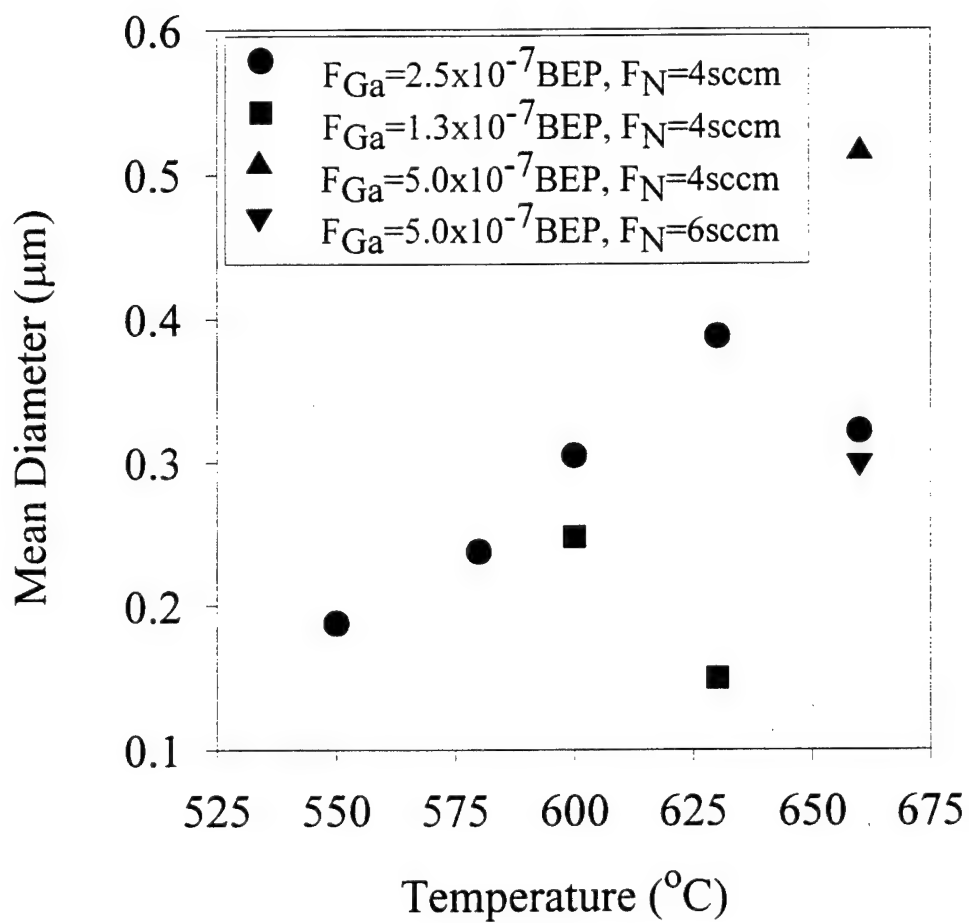


Figure 5.4 GaN nucleation island diameter as a function of growth conditions.

diameters decrease. In contrast, increasing the active nitrogen flux decreases both the maximum island size and the temperature at which that maximum island size occurs. This seems to imply that increasing the substrate temperature for a given set of fluxes favors the growth of critically sized nucleation islands. The critically sized islands are stable to further growth and thus continue to grow, while subcritical islands either incorporate into the growing islands or evaporate. Above the maximum temperature for a given flux, the effect of increasing the temperature is similar to an increase in active nitrogen flux. The process behind this effect is not clearly understood. It is possibly related to an increase in the Ga desorption rate or an increase in the active nitrogen flux through the cracking of metastable nitrogen molecules at the substrate.

If we look at the growth rate of GaN as a function of substrate temperature and Ga flux (Figure 5.5) for a fixed nitrogen flux, we find that for low Ga fluxes there is a linear increase in growth rate with increasing Ga flux. This would be expected if the nitrogen flux present were sufficient to react with all of the available Ga. With further increase in Ga flux, the growth rate levels off, perhaps even declining somewhat, leading eventually to Ga condensation. This implies that the growth rate is limited by the availability of active nitrogen (Ga-rich growth regime). Increasing the flux of active nitrogen in this region increases the growth rate confirming this assertion. The slight decreases in growth rate over the Ga-rich region are most probably related to a transition to a more 2-D growth mode.

Figure 5.6 shows the morphological evolution for samples grown with a Ga flux of 2.5×10^{-7} Torr at substrate temperatures up to the maximum shown in Figure 5.4. The two-fold increase in island diameter yields roughly a factor of four reduction in inter-island defects that need to be smoothed out in subsequent growth. Even with the increase in domain size, the growth is still proceeding in a highly three-dimensional manner leaving a surface with an average roughness greater than 100 nm. This micro-crystallite morphology can be seen more clearly in the height mode image presented in Figure 5.7.

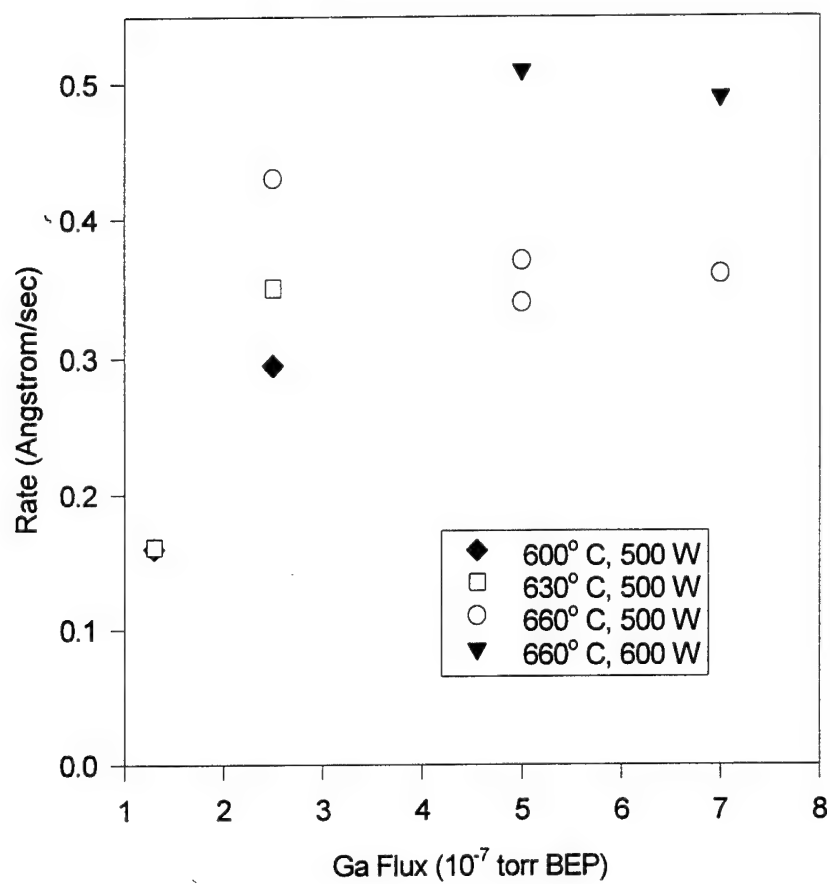


Figure 5.5 Growth rate as a function of Ga flux for several temperatures. Estimated error in growth rate determination is approximately 10%.

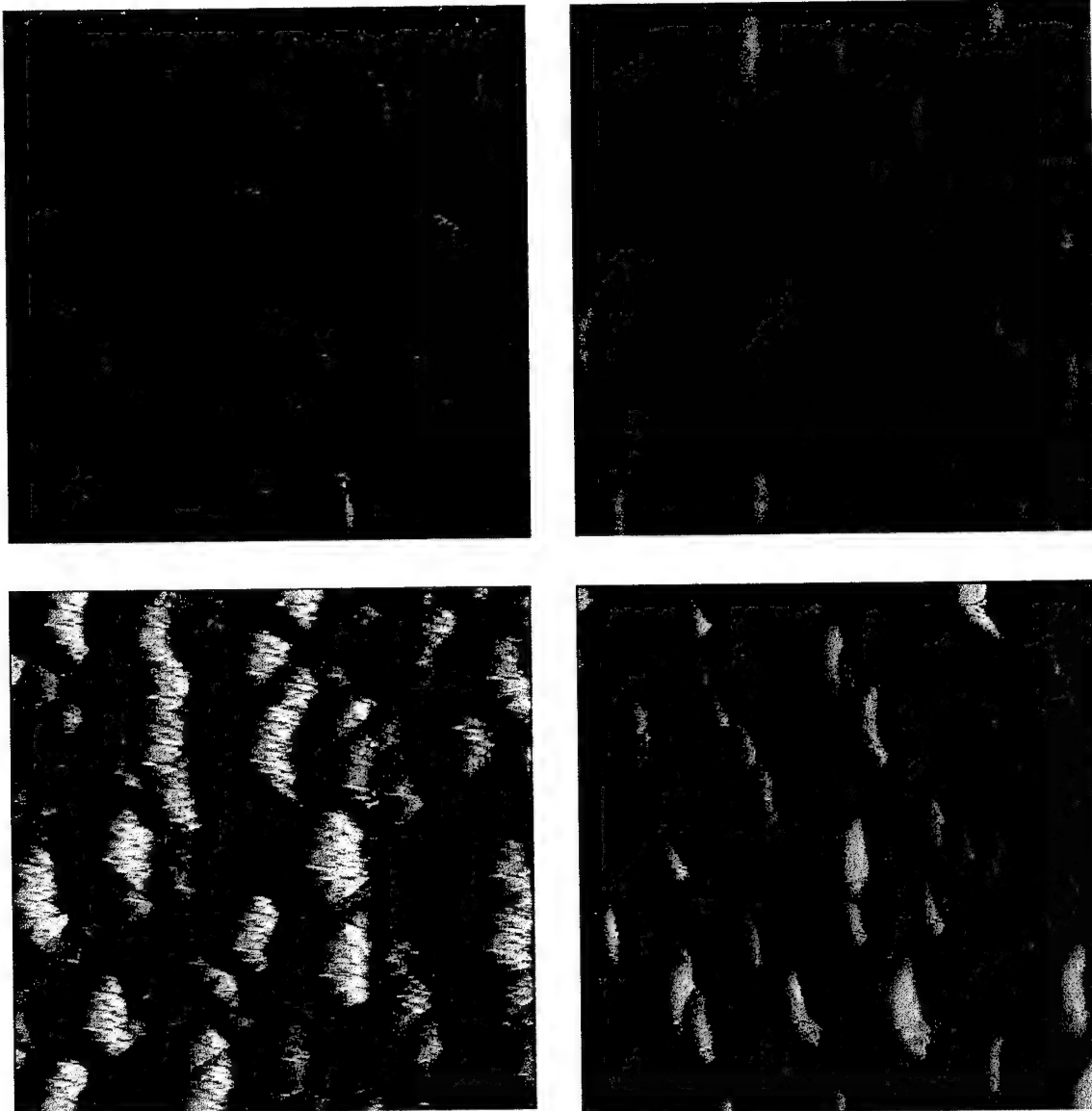


Figure 5.6 (c)Clockwise from top left) AFM micrographs of samples grown under 2.5×10^{-7} torr Ga at substrate temperatures of 550, 580, 600, 630°C and 4 SCCM nitrogen at 500 watts. Measured rms surface roughness was > 100 nm in all cases.

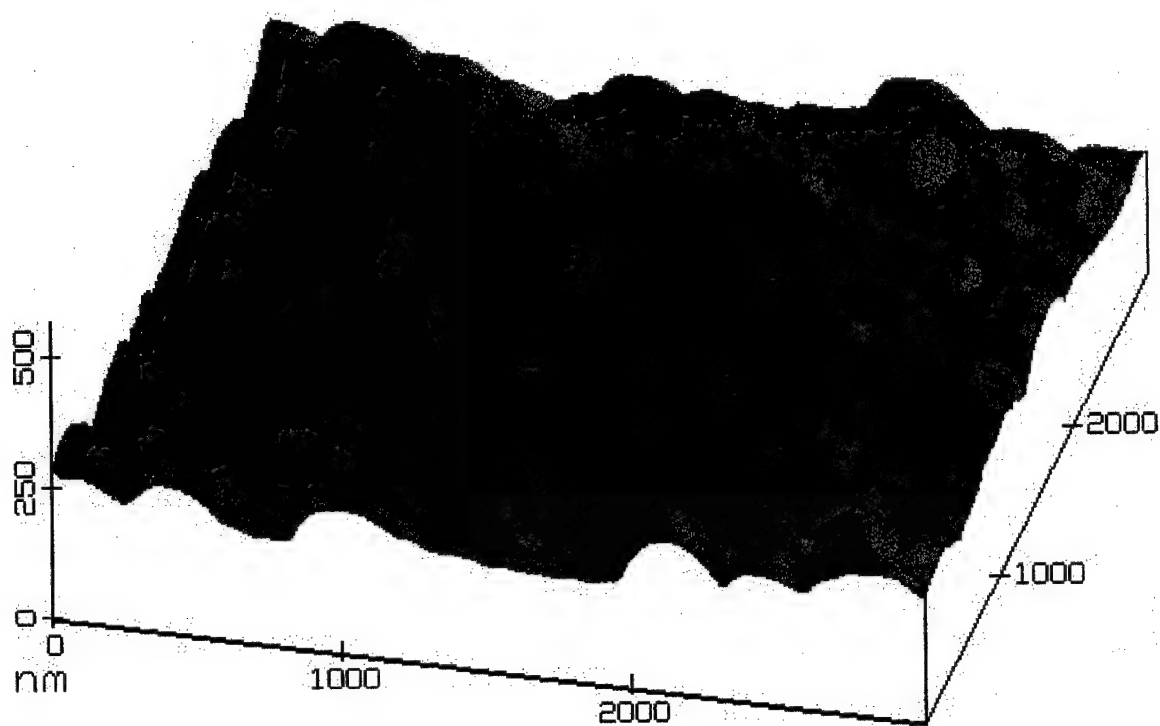


Figure 5.7 Height mode image of one of the samples shown in figure 5.6

Increasing the substrate temperature to 660 °C brings about an interesting change. Figure 5.8a shows an AFM micrograph of a film grown at 660 °C with 2.5×10^{-7} BEP Ga and 4 SCCM N₂ flow at 500 watts. After 3000 Å of growth, the film still exhibits well-defined 3-D growth generating a surface with an average roughness of 150 nm. X-ray measurements showed the film to be single-crystal but with a fairly large full-width-at-half-maximum (FWHM) of 400 arc minutes. The film shown in Figure 5.8b was grown under double the Ga flux used for the sample in Figure 5.8a. The increase in Ga flux brings the film closer to Ga condensation (about 7.0×10^{-7} Torr for this temperature) and produces larger nucleation domains. More importantly, however, is the evident change in growth mode between the two films. The film shown in Figure 5.8b displays the beginnings of a distinctly two-dimensional growth. Nucleation domains are showing signs of coalescence and the average surface roughness has been reduced to about 20 nm. The island tops are relatively flat and have well-defined sub-nanometer steps that correspond to one or two monolayers of growth. X-ray measurements on this film showed a reduction in FWHM to about 120 arc minutes. This morphological change was evident in all films grown in the Ga-rich regime with substrate temperatures higher than 660 °C.

Films grown above 660 °C exhibit signs of the beginnings of 2-D growth. A more standard growth sequence involves an annealing step following a growth interruption. Two samples were grown to investigate the effect of an annealing step on layer morphology. Figure 5.9 shows AFM micrographs of layers grown at 670 °C under 6 SCCM of N₂ at 500 watts and 5.0×10^{-7} BEP Ga. Figure 5.9a again displays flat-topped domains and monolayer terraces. Figure 5.9b shows an identical sample except that growth was interrupted after 100 Å. The film was then annealed under nitrogen at 670 °C for 20 minutes, and growth was resumed. The resulting film displays almost complete coalescence with a surface roughness of about 1.5 nm. X-ray measurements on this film showed a FWHM of roughly 51 arc minutes.

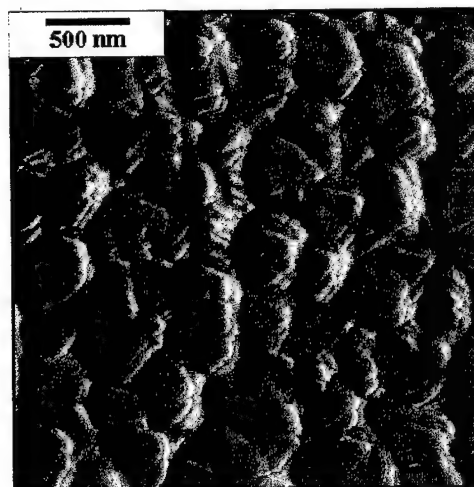
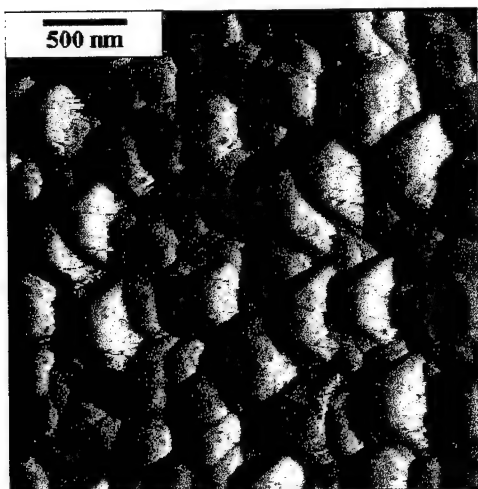


Figure 5.8 AFM micrographs of samples grown at 660°C and 4 SCCM nitrogen at 500 watts. The sample on the left was grown under 2.5×10^{-7} torr Ga while the sample on the right was grown under 5.0×10^{-7} torr Ga.

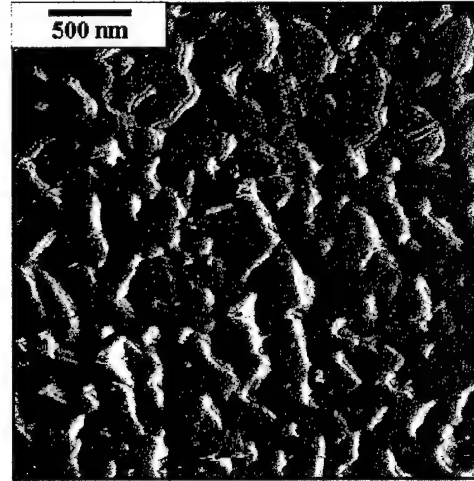
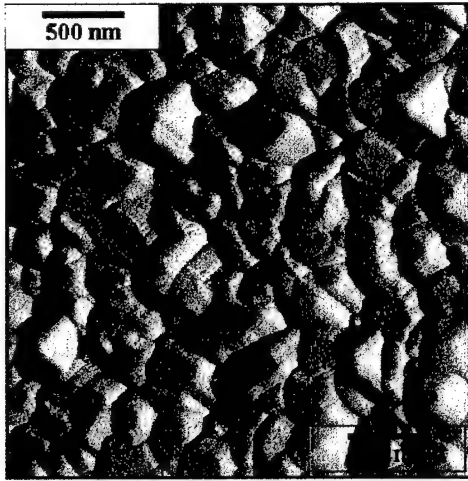


Figure 5.9 AFM micrographs of samples grown at 670°C and 6 SCCM nitrogen at 500 watts. The sample on the left was grown under 5.0×10^{-7} torr Ga while the sample on the right was grown under 7.0×10^{-7} torr.

To briefly summarize then, optimal surface morphologies require growth at substrate temperatures above 660 °C in a Ga-rich flux (nearly at the point of Ga condensation). Annealing the resulting films at or above 660 °C leads to almost complete coalescence of the nucleation domains. By growing films in the Ga-rich regime at substrate temperatures above 660 °C we have found conditions that produce a mean island size of ~0.3 micron (as seen in Figure 5.10). This corresponds to a defect density around $1 \times 10^9 \text{ cm}^{-2}$. Thus, we are poised to begin growth of thicker layers for more complete characterization.

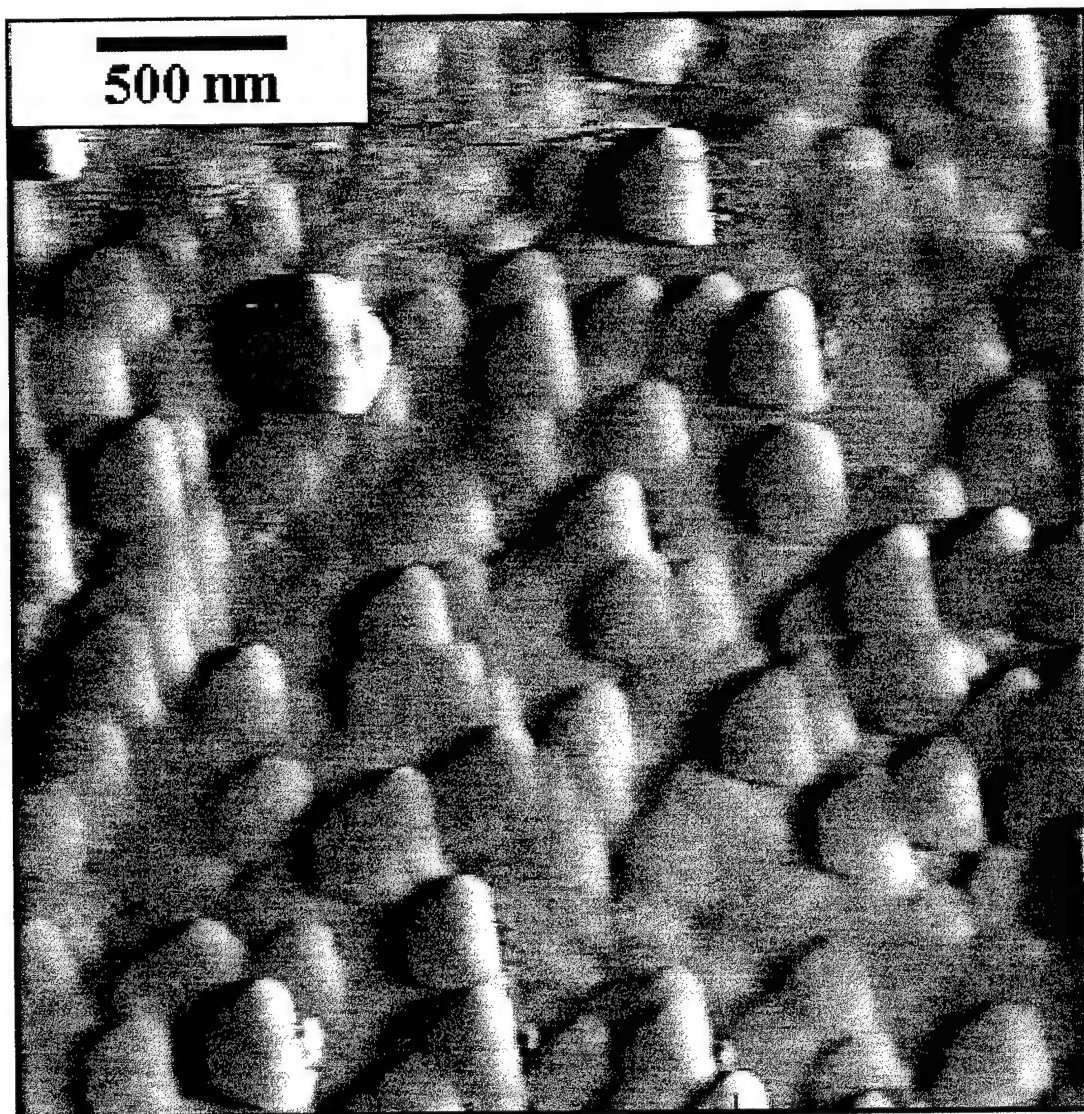


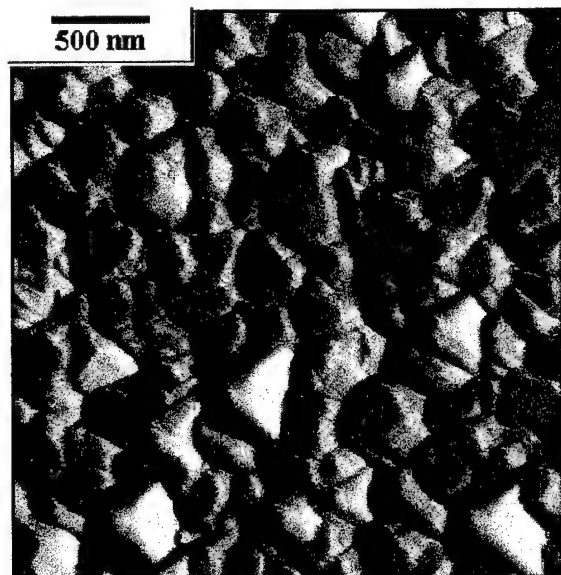
Figure 5.10 Nucleation layer grown at 660°C under 5.0×10^{-7} torr Ga and 6 SCCM nitrogen at 500 watts. The mean diameter of the islands in the sample was 0.3 micron.

6. Growth of GaN

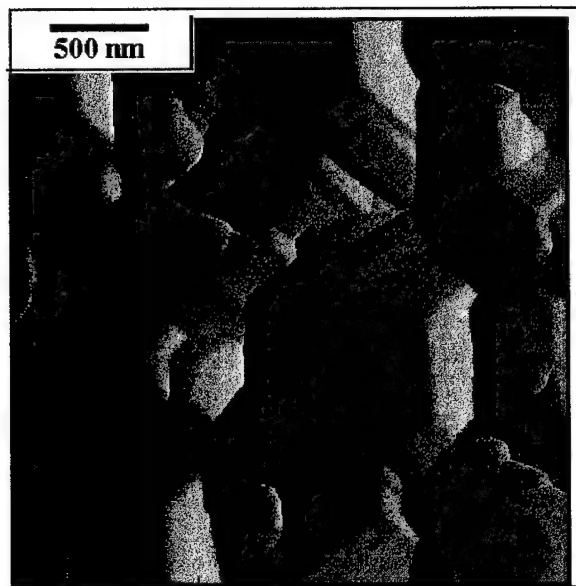
With the established conditions for buffer layer growth as described in the previous section, it was possible to grow films for a more detailed characterization of material properties including PL, AFM, and Hall measurements. This next sequence of layer growths would serve as a baseline to guide subsequent studies using GaN. As part of this study, some samples were grown under a flux of hydrogen (atomic and molecular) to ascertain its effects, if any, on the growth. All films were grown on an optimized GaN buffer layer 100-200Å thick grown under 5.0×10^{-7} Torr Ga at 670 °C and 6 SCCM N₂ at 500 Watts. The buffer layers were then annealed under the nitrogen flux for 10 minutes. The sapphire substrates were prepared as described in Chapter 5 except that several samples were mounted to the molybdenum heater block using molten indium and a molybdenum mask instead of graphite.

6.1 Basic Growth

A series of layers was grown with thicknesses varying from about 0.5 to 1.1 microns. Figures 6.1a and b are AFM micrographs of two such films grown at 670 °C, 6 SCCM and 500 Watts plasma power. Again, this clearly shows the need for Ga-rich growth as the film grown under 7.0×10^{-7} Torr Ga(6.1b) exhibits significantly larger domains than the film grown under 5.0×10^{-7} Torr(6.1a). With further increase in substrate temperature, the surface morphology continues to improve. Figure 6.2 shows two films which were grown at 730 °C under 5.0×10^{-7} Torr Ga, 6 SCCM N₂ and 600 Watts. These films show almost complete coalescence of the nucleation islands with the exception of the pyramidal 'hillocks' evident in Figure 6.2a. The flat areas in both films have an average roughness of ~1.5-2.0 nm and X-ray FWHM of about 4-5 arc minutes. However, increasing the substrate temperature from 670 °C to 730 °C has brought about a



(a)



(b)

Figure 6.1 AFM micrographs of two samples grown at 670°C with 6 SCCM nitrogen at 500 watts.

- a.) Grown under 5.0×10^{-7} torr Ga
- b.) Grown under 7.0×10^{-7} torr Ga

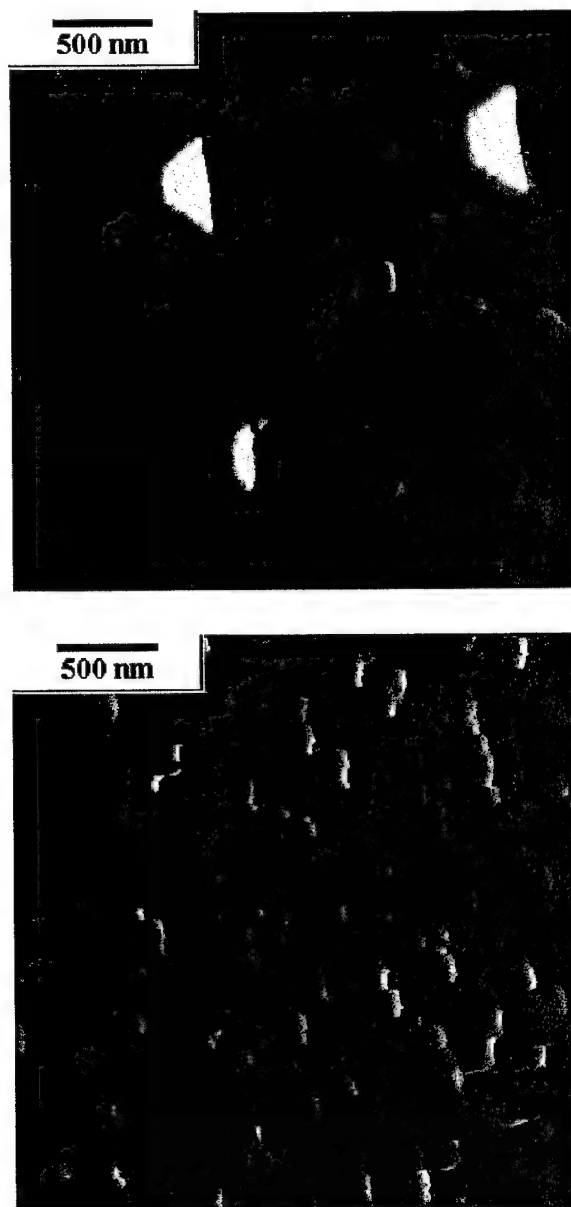


Figure 6.2 AFM micrographs of two samples grown at 730°C and 6 SCCM nitrogen at 600 watts.

(Top) Grown under 5.0×10^{-7} torr Ga

(Bottom) Grown under 7.0×10^{-7} torr Ga

decrease in the overall growth rate, reducing it by. This may be due, in part, to the increasingly 2-D mode of growth exhibited in this film. It may also be due to increased Ga desorption as prior studies have indicated that the Ga desorption rate increases rapidly above 700 °C.

The sample shown in Figure 6.2a has also been examined by Dr. Linda Romano (Xerox PARC) using transmission electron microscopy (TEM). Figure 6.3 is a cross-sectional TEM micrograph of this sample. The dark lines running vertically up from the substrate are the remaining boundaries between nucleation domains. Although most such defects have an edge-type nature, this was not the case in this layer. It appears that sufficient coalescence has removed them, leaving only non-edge type dislocations.

Also, the TEM study identified another type of defect, inversion domain boundaries (IDBs), where the in-plane axes of neighboring domains are effectively reversed relative to one another. A schematic of the IDB and its possible relation to substrate surface inhomogeneities is shown in Figure 6.4.

In sharp contrast to GaN grown in other labs by MBE, the IDBs in our sample are estimated to fill less than 10% of the sample volume. Typical MBE grown GaN films exhibit defect levels around 50% [30].

By counting the dislocations which intersect the surface, as seen on Figure 6.3, it is possible to estimate the dislocation density. For this film, the density works out to roughly $2 \times 10^9 \text{ cm}^{-2}$ which compares favorably with the best reported by any other growth technique (MOCVD: $\sim 7 \times 10^8 \text{ cm}^{-2}$). This also compares quite well with the density of $1 \times 10^9 \text{ cm}^{-2}$ estimated from nucleation island size and resultant structure observed in AFM from our AFM study on layers grown with similar nucleation conditions. Note, however, that even though this defect density is comparable to the lowest reported, it is still five orders-of-magnitude higher than would be tolerable in other semiconductor systems, like GaAs.



Figure 6.3 Dark field TEM image of sample 9571 (AFM image shown in figure 6.2a). 'ID' denotes an Inversion Domain. The remaining defects are non-edge type dislocations. (Photo courtesy of Dr. Linda Romano, Xerox PARC)

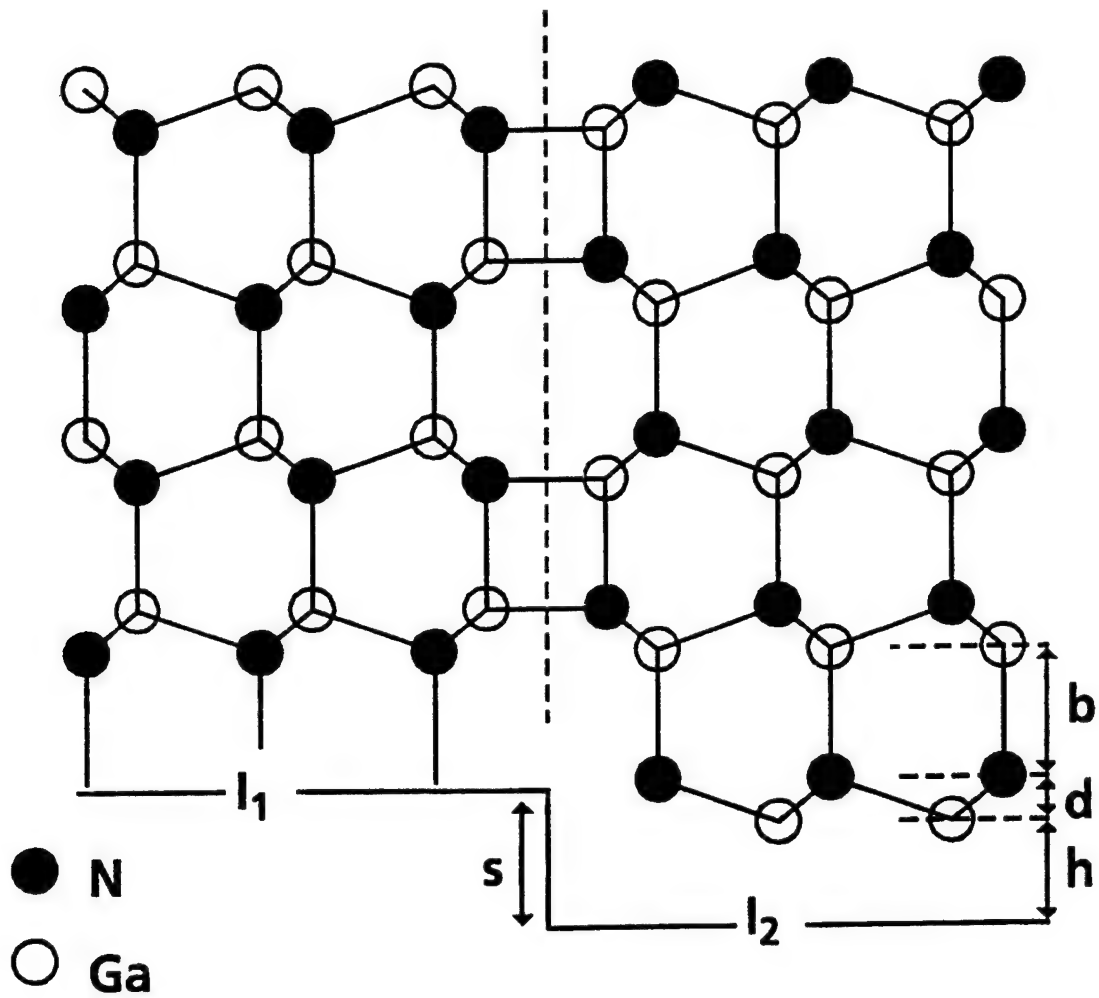


Figure 6.4 Schematic of Inversion Domain structure. ('S' represents a substrate step or defect which may be responsible for creating this type of defect.) (Drawing courtesy of Dr. Linda Romano, Xerox, PARC)

Photoluminescence was performed at liquid helium temperature by Prof. Giles in her laboratory on several samples prepared in this growth run. The samples exhibited band-edge emission at about 365 nm (3.47 eV). This is the so-called I_1 peak, and is commonly ascribed to an exciton bound to a nitrogen vacancy. However, this assignment has not been clearly proven. The I_1 peak is, however, typically observed for material grown under Ga-rich (nitrogen deficient) conditions. All samples grown at or below 660 °C displayed an additional broad luminescence band centered at about 560 nm (2.2 eV), as seen in Figure 6.5a. While the origin of this yellow luminescence is still controversial [31], it is commonly attributed to deep states in the bandgap involving impurities or native defects [32]. There is a growing belief in the community that this luminescence is associated with edge-type dislocations occurring at coalescence boundaries. Interestingly, the layers grown at 730 °C did not exhibit a detectable level of this yellow luminescence, as seen in Figure 6.5b. This may indicate that temperatures above 660 °C are necessary for high quality material. In low temperature PL measurements, the layers grown at 730 °C exhibited primarily near-bandedge luminescence related to excitonic transition, again suggesting an improvement in at least those aspects of layer quality probed by PL.

The yellow luminescence seen in the PL data can also be seen, at room temperature, with the aid of an optical microscope with an UV light source. Samples exhibiting the yellow PL band could be seen to shine bright yellow under UV excitation. With the aid of an optical microscope, it was possible to perform UV fluorescence microscopy (UVFL) to examine the spatial distribution of the yellow emission. Figure 6.6 shows one such micrograph. One can clearly see individual microcrystallites resolvable as 0.5-0.3 micron in diameter. It is apparent that the luminescence is not uniform across the sample and varies from one microcrystallite to another. Some isolated microcrystallites exhibited blue luminescence. Occasional samples exhibited weak PL centered near 400 nm or 440 nm when measured using the

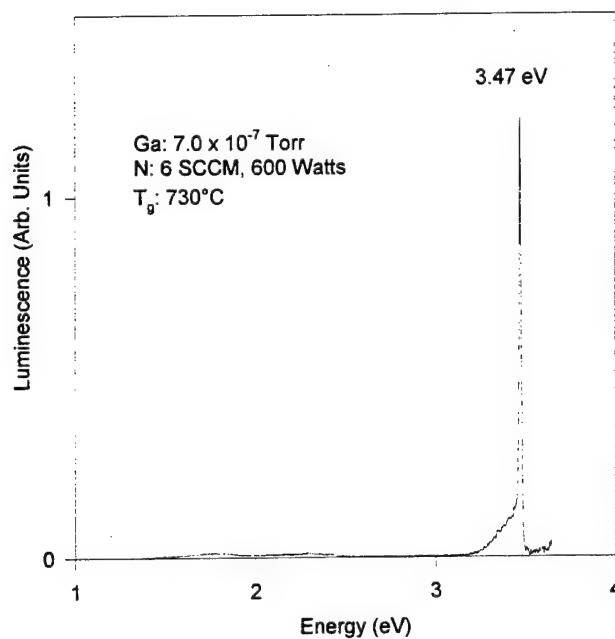
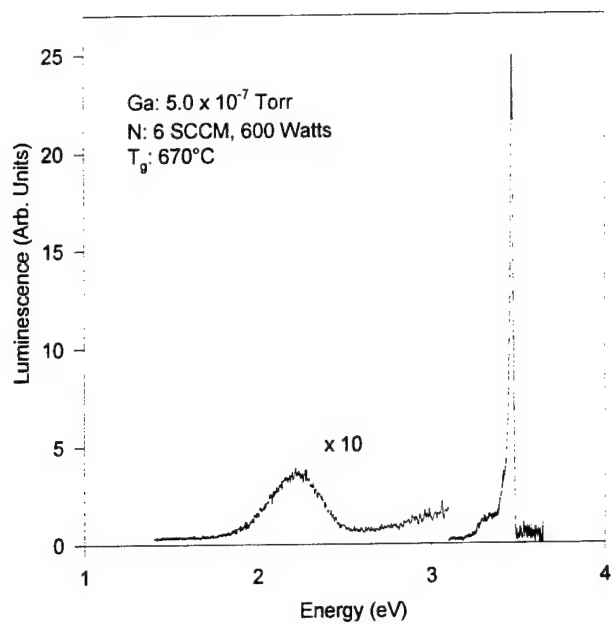


Figure 6.5 Photoluminescence spectra for two samples grown at 660°C (top) and 730°C (bottom). Note suppression of the 2.2 eV 'yellow' luminescence in the higher temperature growth.

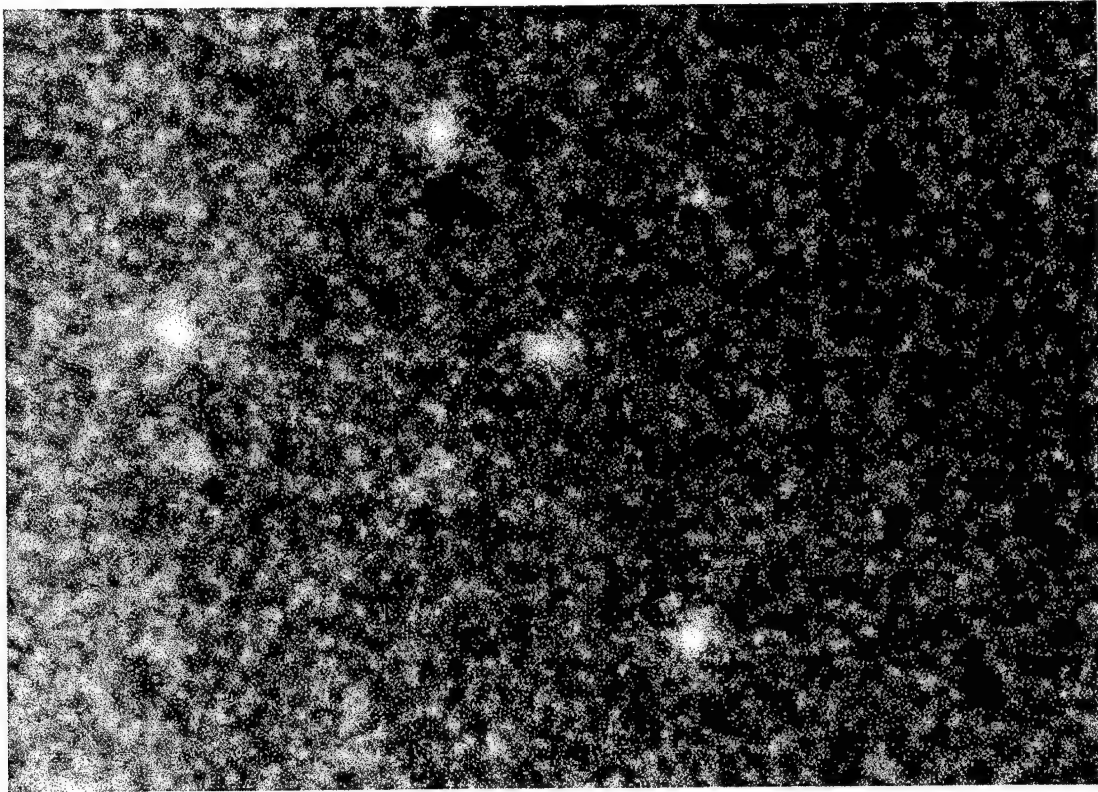


Figure 6.6 Ultraviolet Fluorescence micrograph of 'yellow luminescence' seen in PL spectra.

standard PL setup. The UVFL result indicates that this may actually originate from localized impurities or defects. The samples grown at 730 °C did not exhibit any detectable yellow fluorescence at room temperature.

Hall measurements were performed on these samples at room temperature. A clear trend emerged in these measurements. For samples grown at 730 °C, we observed the high n-type carrier concentration ($6.3 \times 10^{17} - 2.5 \times 10^{18} \text{ cm}^{-3}$) reported for most samples grown by any technique. Mobilities ranged from 4.9 cm^2/Vs to 469.2 cm^2/Vs , with the lowest values coming predominantly from the higher temperature growths. Of interest is one measurement indicating a mobility of 4,000 cm^2/Vs at low carrier concentration. This, of course, needs to be carefully re-evaluated and a temperature dependent study performed to determine compensation ratios.

This n-type conduction is often attributed to nitrogen vacancies, as it is believed that most higher temperature growths take place under conditions conducive to vacancy formation [33]. An alternative theory is that the n-type carrier concentration is related to defects resulting from the decomposition of the GaN during typical growth conditions [34]. This would occur where the growth rate starts decreasing with temperature with the attendant increase in Ga desorption rate. Rubin, et al. [35], observed a dramatic decrease in n-type carrier concentration upon going to lower growth temperatures, with mobilities as high as 1200 cm^2/Vs for a carrier concentration of 10^{14} cm^{-3} . In addition, they observed as-grown p-type conduction, possibly related to a reduction of the n-type background due to this unknown defect.

We also observed a dramatic decrease in n-type carrier concentration upon going to the low temperature growths. For samples grown at 660 °C or less, carrier concentrations were both n- and p-type, ranging from 9×10^{12} to $4 \times 10^{15} \text{ cm}^{-3}$. The mobilities covered a wide range, probably indicating various degrees of compensation and possibly mixed conduction in some cases.

A future study may be to use higher temperatures to obtain structural perfection, followed by lower temperature growth to eliminate (or minimize) this n-type defect.

6.2 Effects of Hydrogen on the Growth of GaN

It is well known that hydrogen easily bonds to the surface of other semiconductor systems, such as silicon or germanium [36]. There is also evidence that atomic hydrogen alters the growth kinetics in GaAs [37,38]. For several samples, hydrogen was incorporated into the growth flux during GaN growth. H_2 passed through the thermal cracker described earlier produced a low flux of atomic hydrogen. The unfilled symbols in Figure 6.7 show the change in growth associated with the introduction of hydrogen. As much as a factor of two increase in growth rate was seen for all samples grown under Ga-rich conditions with hydrogen, when compared to their hydrogen-free counterparts. This is perhaps the most intriguing finding of this study. Based on this, it is likely that hydrogen is playing an important role in the growth kinetics of MOCVD, MOMBE, and in GSMBE using ammonia. A detailed study of this phenomenon is currently underway in Prof. Myers' research group, funded by the Office of Naval Research from a proposal based on the research described in this thesis.

Hydrogen incorporating directly into the film cannot account for the increase on growth rate. Figure 6.8 shows two films grown under identical conditions except that one was grown with an additional 1×10^{-6} Torr hydrogen in the vacuum chamber. It is readily seen that the film grown with hydrogen has a much smaller domain size than the hydrogen-free film. It was shown earlier that both a decrease in domain size and an increase in growth rate, in the Ga-rich regime, can be explained by a shift to a more nitrogen sufficient growth. This implies that the presence of hydrogen effects the growth kinetics, effectively increasing the active nitrogen flux. It is possible that the hydrogen bonds loosely to the growing surface where it attracts incident nitrogen atoms. The

Growth Rate vs. Ga Flux

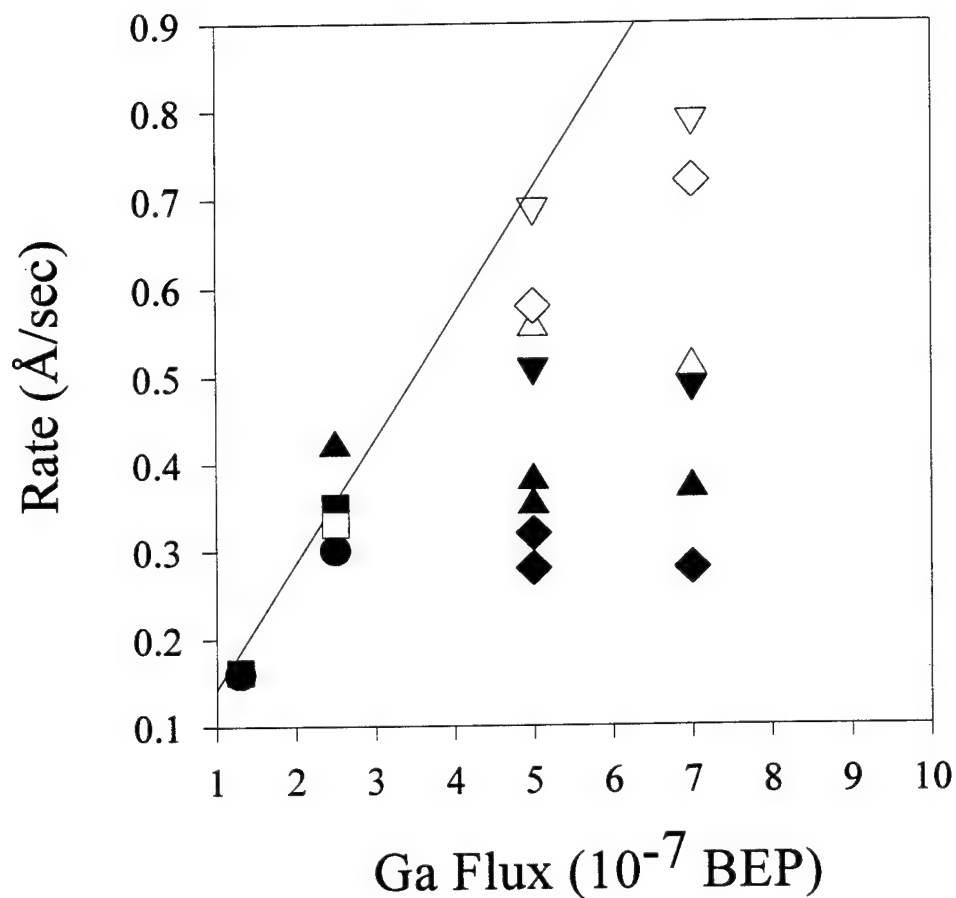


Figure 6.7 Graph of growth rate versus Ga flux for various growth conditions. Unfilled symbols correspond to growth conditions of similar filled symbols but with the addition of atomic hydrogen to the growth flux.

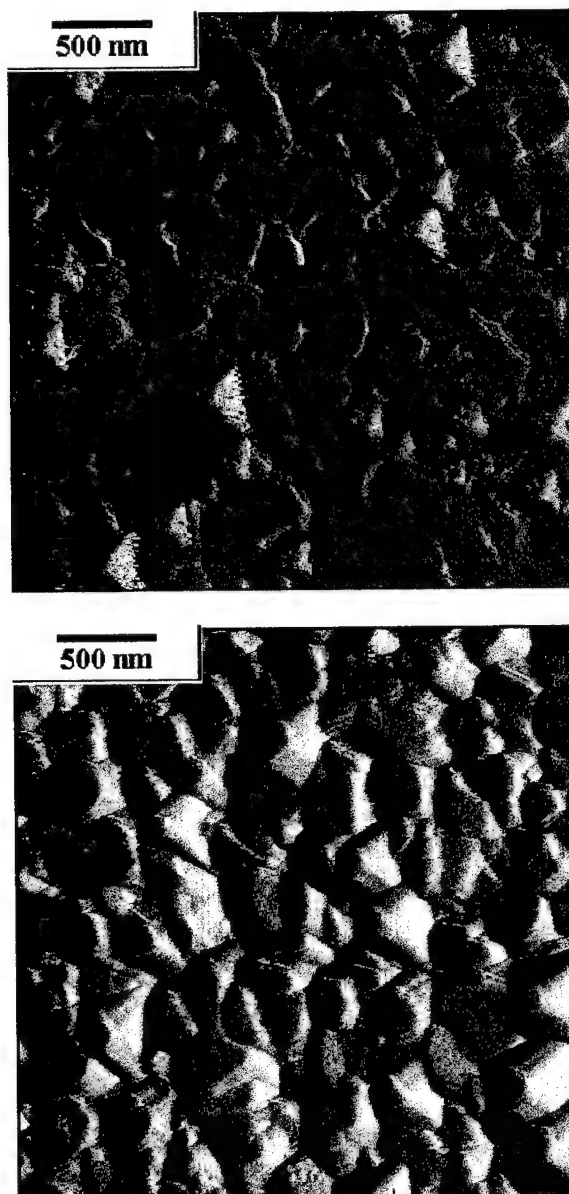


Figure 6.7 AFM micrographs of two samples grown at 670°C under 5.0×10^{-7} torr Ga and 6 SCCM nitrogen at 500 watts.
a.) Without hydrogen
b.) With 1.0×10^{-6} torr added to growth flux through thermal cracker. Note smaller domain size similar to increasing nitrogen flux.

nitrogen atoms attach to this hydrogen layer thereby increasing the nitrogen residence time. This allows adsorbed Ga atoms more time to diffuse to within an interaction distance to bond with the nitrogen. Doubling the hydrogen flux to 2.0×10^{-6} Torr resulted in no noticeable difference in growth rate, implying that the reaction does not depend on the incident flux of hydrogen, just on its presence.

Structural and electrical properties were indistinguishable between films grown with and without hydrogen. The best GaN films grown with hydrogen exhibited x-ray diffraction FWHM between 4 and 5 arc minutes. Hall measurements suggested a possible factor of two improvement in carrier concentration, and an order-of-magnitude improvement in mobility, between similar samples.

Figure 6.9 shows PL results for two films grown with hydrogen. Again the low temperature growth exhibited a yellow luminescence band while the high temperature growths exhibited mainly band-edge luminescence. Figure 6.9b shows the 3.47 eV bound-exciton peak and the absence of PL related to deep levels, indicating that growth under hydrogen does not degrade layer quality.

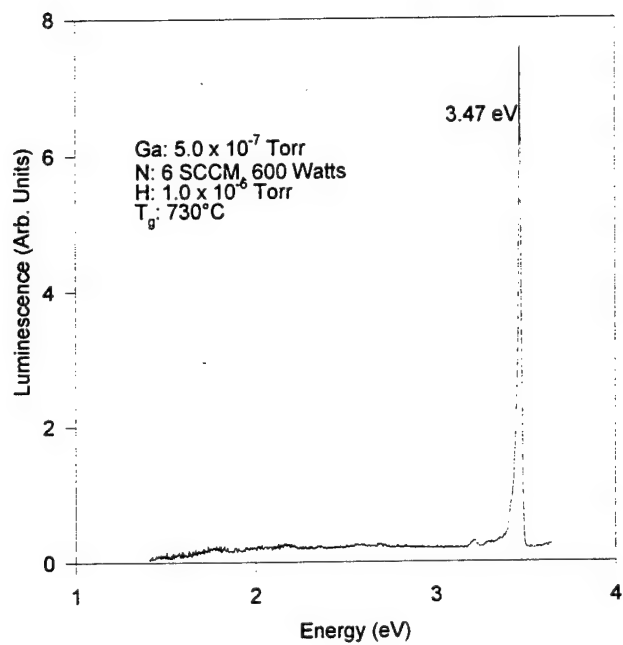
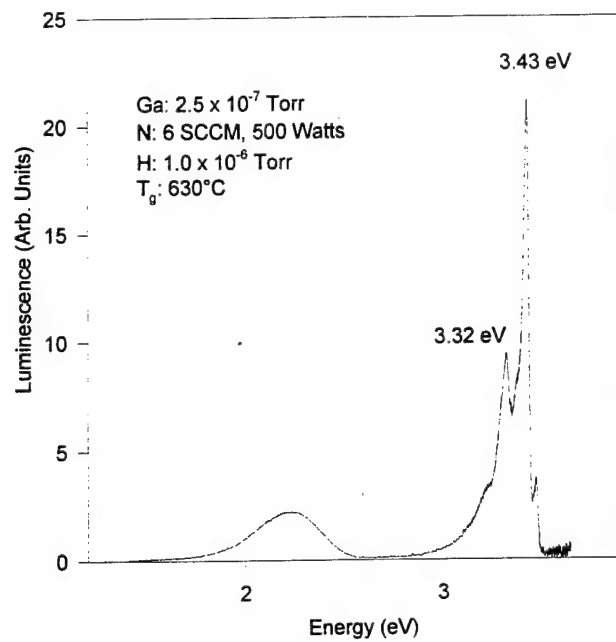


Figure 6.9 PL spectra for two samples grown with atomic hydrogen in the growth flux. Spectra are virtually unchanged compared to similar non-hydrogen growths.

7. Summary and Future Areas for Research

GaN is an interesting, challenging and technologically important semiconductor system. Its potential applications range from acid-resistant coatings to optoelectronic devices operating in the blue and near-UV wavelengths. For this reason, considerable effort has been, and is being, expended by this lab, and by others around the world, toward growing high quality material and understanding its properties. In recent years, numerous advances in the growth and processing of GaN have sparked new interest in its potential. Newer growth techniques have improved the crystallinity and helped demonstrate p-type material growth. The use of buffer layers for growth on sapphire substrates has significantly improved layer quality, both electrically and mechanically. The upcoming availability of better lattice-matched substrates and, possibly, GaN substrates for homoepitaxial growth are sure to bring about even greater improvements.

The four major goals of my research were achieved. Characterization of the RF radical source was conducted using optical emission and mass spectroscopies. It was shown that the output flux consists of atomic and molecular nitrogen, and ions of each in varying concentrations. For a given source operating power, the percent of the flux converted to atomic nitrogen is shown to decrease with increasing gas flow into the source. When converted to actual fluence, however, the atomic nitrogen flux increases with increasing flow and power. Ion fluxes were shown to be low (less than 0.03% of the total output) for this source when using an aperture plate with small (<0.5 mm diameter) exit holes. Larger exit holes exhibit higher fluxes of high energy ions. This is likely due to a decrease in the recombination of ions in the boundary layer of the exit holes.

As a part of this research effort, thirty-two GaN films have been grown to date in the WVU Molecular Beam Epitaxy lab. The primary goal of these growths was to study the initial nucleation of GaN on sapphire. Nucleation layers were grown at substrate

temperatures ranging from 550°C to 670 °C, under Ga fluxes ranging from $1.3\text{-}5.0 \times 10^{-7}$ Torr. Active nitrogen was supplied by a RF nitrogen source. Examination of layer surface morphologies by atomic force microscopy showed significant improvement in layer quality when growths were conducted at higher temperatures under a Ga-rich growth flux. In fact, the highest layer quality occurred at Ga fluxes just short of where Ga begins to condense out onto the sample surface. Growths under these conditions displayed the beginnings of a change in growth mode from 3-D micro-crystallites to a flatter, more 2-D growth. These growth conditions were also shown to promote larger nucleation islands, which in turn lowers the final density of domain-edge related crystal defects. Conditions were reported which resulted in $2 \times 10^9 \text{ cm}^{-2}$ defect densities, comparable to the $7 \times 10^8 \text{ cm}^{-2}$ commonly considered to be state-of-the-art. Annealing of the nucleation layers was shown to promote a high degree of domain coalescence, resulting in films with average surface roughness less than 5 nm.

Thicker layers ($<1 \text{ }\mu\text{m}$) were grown for some of the higher temperature, higher quality conditions. Characterization by photoluminescence showed that growth at lower temperatures led to the phenomenon of "yellow luminescence", commonly attributed to poor structural quality. This luminescence was greatly suppressed in layers grown above 670 °C. TEM measurements confirmed the low density of large-scale crystalline defects and demonstrated that those remaining were primarily non-edge type dislocations, giving hope of further reductions in density with two dimensional growth.

Several films were grown with atomic hydrogen added to the growth flux. This resulted in growth rate enhancements of as much as a factor of two. The hydrogen is most likely altering the growth kinetics, increasing the residence time of nitrogen on the sample surface. Inclusion of hydrogen was shown to have no discernible effect on material properties under the conditions utilized here.

Work is already in progress to further the study reported here. Much interest has been expressed in the effects of hydrogen on growth kinetics. Further work needs to be

done with growths under a wider range of hydrogen fluxes to determine if there is a limit to the possible benefits of this technique. Growth techniques need to be further refined to promote the early onset of 2-D growth and further reduce the level of intrinsic crystal defects. Further material characterization also needs to be conducted to determine what growth conditions are necessary, not only for high structural quality but high electrical and optical quality as well. In addition, recent SIMS studies of two samples indicated a potentially disturbing fact; at least those samples, and quite likely most of the others, are heavily contaminated with boron (see figure 7.1). The contamination most likely stems from the breakdown of the pyrolitic boron nitride crucible in the nitrogen source. A new crucible has been installed and future growths will hopefully determine the effect of the presence of boron. In the end, it may prove that boron inclusion is necessary for high quality layer growth.

Thus, the research presented in this thesis represents a solid baseline for future efforts in understanding the growth and properties of GaN.

GNSA-9567

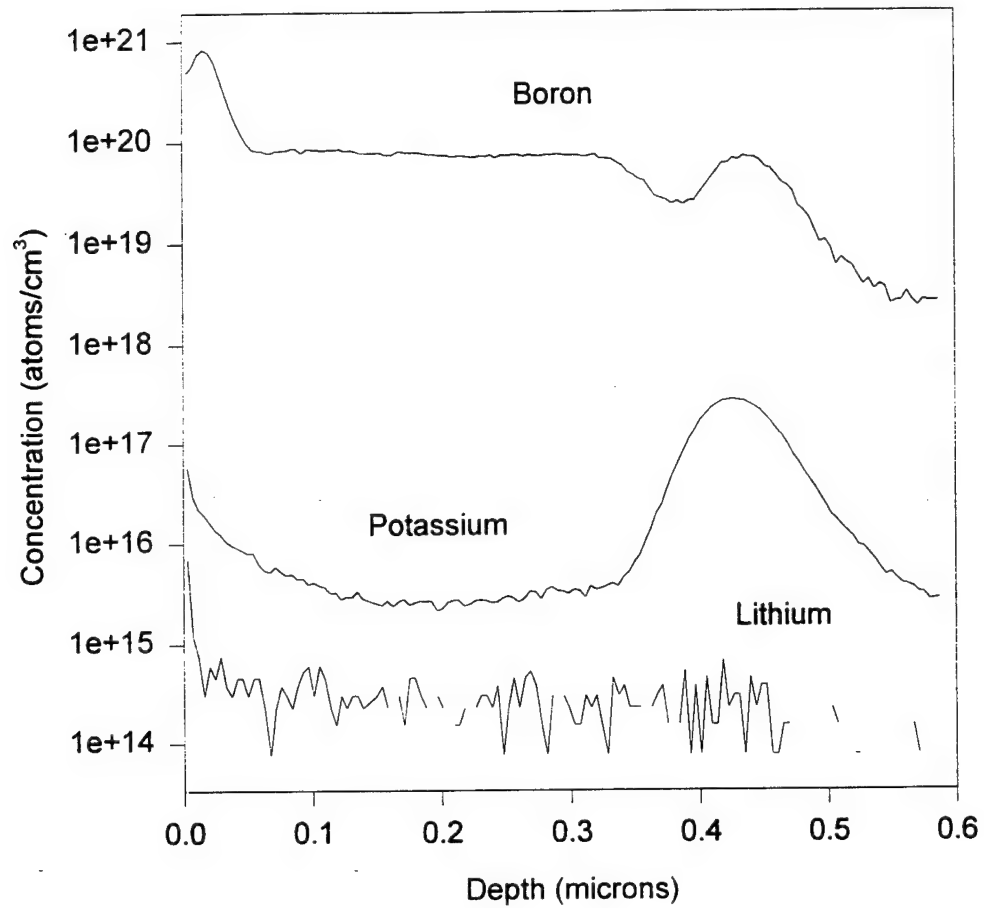


Figure 7.1 SIMS spectrum for sample 9567 indicating a very high level of boron inclusion. Similar results were obtained for sample 9569. This may indicate boron inclusion in all samples grown to date.

References

- [1] H. Morkoç, S. Strite, G.B. Gao, M.E. Lin, B. Sverdlov and M. Burns, *J. Appl. Phys.* **76**, 1363 (1994).
- [2] Z. Yu, PhD Dissertation, Physics Dept., WVU (1996).
- [3] See for example: *Mat. Res. Soc. Proc.* Vol. 283 (1993).
- [4] W. Götz, N.M. Johnson, J. Walker, D.P. Bour, H. Amano and I. Akasaki, *Appl. Phys. Lett.* **67**, 2666 (1995).
- [5] M.S. Brandt, N.M. Johnson, R.J. Molnar, R. Singh and T.D. Moustakas, *Appl. Phys. Lett.* **64**, 2264 (1994).
- [6] H. Amano, M. Kito, K. Hiramatsu and J. Akasaki, *Jpn. J. Appl. Phys.* **28**, L2112 (1989).
- [7] S. Nakamura, T. Mukai, M. Senoh and N. Iwasa, *Jpn. J. Appl. Phys.* **31**, L139 (1992).
- [8] S. Nakamura, T. Mukai and M. Senoh, *Jpn. J. Appl. Phys.* **30**, L1998 (1991).
- [9] R.C. Powell, N.E. Lee, Y.W. Kim and J.E. Greene, *J. Appl. Phys.* **73**, 189 (1993).
- [10] R.J. Molnar and T.D. Moustakas, *J. Appl. Phys.* **76**, 4587 (1994).
- [11] Z. Yang, L. Li and W. Wang, *Appl. Phys. Lett.* **67**, 1686 (1995).
- [12] M. Asif Khan, T.D. Olson, J.N. Kuznia, W.E. Carlos and J.A. Freitas, Jr. **74**, 5901 (1993).
- [13] S. Nakamura, T. Mukai and M. Senoh, *J. Appl. Phys.* **71**, 5543 (1992).
- [14] S. Strite and H. Morkoç, *J. Vac. Sci. Technol. B* **10**, 1237 (1992).

- [15] J.S. Hwang, A.V. Kuznetsov, S.S. Lee, H.S. Kim, J.G. Choi and P.J. Chong
Journal of Crystal Growth **142**, 5 (1994).
- [16] J.N. Kuznia, M. Asif Khan and D.T. Olson, J. Appl. Phys. **73**, 4700 (1993).
- [17] A.Y. Cho and J.R. Arthur, Progress in Solid-State Chemistry **10**, 157 (1975).
- [18] J.A. Venables, G.D.T. Spiller and M. Hanbücken, Rep. Prog. Phys. **47**, 399
(1984).
- [19] B.A. Joyce, Rep. Prog. Phys. **37**, 363 (1974).
- [20] L.J. Van der Pauw, Philips Res. Rep. **13**, 1 (1958).
- [21] G. Binnig, C.F. Quate and Ch. Gerber, Phys. Rev. Lett. **56**, 930 (1986).
- [22] D. Rugar and P. Hansma, Physics Today, 23 (October 1990).
- [23] See also: S. Kumar, MS Thesis, Dept. of Chem. Eng., 1996.
- [24] R.P. Vaudo, J.W. Cook, Jr. and J.F. Schetzina, J. Crystal Growth **138**, 430 (1994).
- [25] W.E. Hoke, P.J. Lemonias and D.G. Weir, J. Crystal Growth **111**, 1024 (1991).
- [26] R.P. Vaudo, J.W. Cook, Jr. and J.F. Schetzina, J. Vac. Sci. Technol B **12**, 1232
(1994).
- [27] R.P. Vaudo, Z. Yu, J.W. Cook, Jr. and J.F. Schetzina, Optics Letters **18**, 1843
(1993).
- [28] J.M. Van Hove, G.J. Cosimini, E. Nelson, A.M. Wowchak and P.P. Chow,
Journal of Crystal Growth **150**, 908 (1995).
- [29] T.D. Moustakas, T. Lei and R.J. Molnar, Physica B, **185**, 36 (1993).

- [30] L.T. Romano, J.E. Northrup and M.A. O'Keefe, Appl. Phys. Lett. **69**, 2394 (1996).
- [31] P. Kung, X. Zhang, D. Walker, A. Saxler, J. Piotrowski, A. Rogalski and M. Razeghi, Appl. Phys. Lett. **67**, 3792 (1995).
- [32] R. Singh, R.J. Molnar, M.S. Ünlü and T.D. Moustakas, Appl. Phys. Lett. **64**, 336 (1994).
- [33] J. Neugebauer and C.G. Van de Walle, Phys. Rev. B **50**, 8067 (1994).
- [34] N. Newman, J. Ross and M. Rubin, Appl. Phys. Lett. **62**, 1242 (1993).
- [35] M. Rubin, N. Newman, J.S. Chan, T.C. Fu and J.T. Ross, Appl. Phys. Lett. **64**, 64 (1994).
- [36] J.A. Schaefer, Physica B **170**, 45 (1991).
- [37] Y.J. Chun, Y. Okada and M. Kawabe, Jpn. J. Appl. Phys. **32**, L1085 (1993).

Appendix A: List of growth conditions for GaN samples

Sample Number	Growth Temp.	Pga (10e-7 Torr)	N (SCCM)	Power (Watts)	Sub. etch	H(e-6)	N PR	mn t	rate (Å/s)	Gro (hrs)	Thick (μm)	Buf (Å)	BPga (e-7)	BN2 (SCCM)	Anneal (min)	BTm
GNSA-9544	630	2.5	4	500	no	no	no	Gra	0.37	6	0.8	no				
GNSA-9545	580	2.5	4	500	yes	no	no	Gra	0.32	6	0.7	no				
GNSA-9546	600	2.5	4	500	yes	no	no	Gra	0.30	6	0.65	no				
GNSA-9547	550	2.5	4	500	yes	no	no	Gra	0.30	6	0.65	no				
GNSA-9548	630	2.5	4	500	yes	no	no	Gra	0.35	6	0.75	no				
GNSA-9549	600	5	4	500	yes	no	no	Gra	0.00	6		no				
GNSA-9550	630	1.3	4	500	yes	no	no	Gra	0.00	6		no				
GNSA-9551	630	5	4	500	yes	no	no	Gra	0.16	6	0.35	no				
GNSA-9552	600	1.3	4	500	yes	no	no	Gra	0.37	3	0.4	no				
GNSA-9553	660	5	4	500	yes	no	no	Gra	0.35	6	0.75	no				
GNSA-9554	660	2.5	4	500	yes	no	no	Gra	0.32	3	0.35	no				
GNSA-9555	630	2.5	4	500	yes	yes	no	Gra	0.56	3	0.6	no				
GNSA-9556	630	2.5	4	500	yes	no	no	Gra	0.00	3		no				
GNSA-9557	630	5	6	500	yes	no	no	Gra	0.00	3		50	2.5			450
GNSA-9558	630	2.5	4	500	yes	no	no	Gra	0.32	3	0.35	no				
GNSA-9559	660	5	6	500	yes	no	no	Gra	0.00	0.05						
GNSA-9560	660	5	6	500	yes	no	no	Gra	0.27	3	0.29	100	5			
GNSA-9561	670	5	6	500	yes	no	no	Gra	0.56	3	0.6	100	5			
GNSA-9562	670	5	6	500	yes	no	no	Gra	0.37	3	0.4	100	5			
GNSA-9563	670	7	6	500	yes	no	no	Gra	0.51	3	0.55	200	7			
GNSA-9564	670	7	6	500	yes	2	no	Gra	0.56	3	0.6	200	5			
GNSA-9565	670	7	6	500	yes	2	no	Gra	0.02	6	0.045	200	5			
GNSA-9566	670	2.5	6	0	yes	no	no	Gra	0.51	3	0.55	200	5	6	10	670
GNSA-9567	670	5	6	600	yes	no	no	Gra	0.79	3	0.85	200	5	6	10	670
GNSA-9568	670	7	6	600	yes	1	no	Gra	0.69	3	0.75	200	5	6	10	670
GNSA-9569	670	5	6	600	yes	1	no	Gra	0.32	3	0.35	200	5	6	10	670
GNSA-9570	670	5	6	600	yes	1-H2	no	Gra	0.32	3	0.35	200	5	6	10	670
GNSA-9571	730	5	6	600	yes	no	no	In	0.32	6	0.7	200	5	6	10	670

Appendix A: List of growth conditions for GaN samples

Sample Number	Growth Temp.	Pga (10e-7 Torr)	N (SCCM)	Power (Watts)	Sub. etch	H(e-6)	N PR	mn t	rate (Å/s)	Gro (hrs)	Thick (µm)	Buf (Å)	BPga (e-7)	BN2 (SCCM)	Anneal (min)	BTm
GNSA-9572	730	7	6	600	yes	no	no	In	0.28	6	0.6	200	5	6	10	670
GNSA-9573	730	5	6	600	yes	1	no	In	0.56	6	1.2	200	5	6	10	670
GNSA-9574	730	5	6	600	yes	no	no	In	0.25	12	1.1	200	5	6	10	670
GNSA-9575	730	5	6	600	y	no			0.28	6	0.6	200	5	6	10	670
GNSA-9611	730	7	6	600	y	1	no	In	0.00	6		200	5	6	10	
GNSA-9664	730	5	6	600	yes	no	no	In	0.56	7	1.4	200	5	6	10	730
GNSA-9665	730	5	6	600	yes	no	no	In	0.44	6	0.95	200	5	6	10	730
GNSA-9666	730	5	6	600	yes	no	no	In	0.00	6		200	5	6	10	730
GNSA-9667	730	5	6	600	yes	no	no	In	0.61	3	0.66	200	5	6	10	730
GNSA-9668	730	7	6	600	yes	no	no	In	0.39	3	0.42	200	7	6	10	730
GNSA-9669	730	9	6	600	yes	no	no	In	0.37	3	0.4	200	9	6	10	730
GNSA-9670	730	9	6	600	yes	1	no	In	1.11	3	1.2	200	5	6	10	730

Appendix A: List of growth conditions for GaN samples

Sample Number	Carrier Concentration	Mobility	Notes
GNSA-9544			
GNSA-9545			
GNSA-9546	-4.30E + 15	280.7	
GNSA-9547			
GNSA-9548			
GNSA-9549			
GNSA-9550			
GNSA-9551			
GNSA-9552			
GNSA-9553			
GNSA-9554			
GNSA-9555			
GNSA-9556			
GNSA-9557			
GNSA-9558	9.20E + 12	46.4	
GNSA-9559	-2.10E + 15	31.8	
GNSA-9560			
GNSA-9561	4.90E + 15	10.1	Buffer layer annealed 20 minutes under N2
GNSA-9562			Same anneal
GNSA-9563			same anneal
GNSA-9564	-1.10E + 15	469.2	Annealed 10 minutes
GNSA-9565		428.8	Same anneal Buffer grown at 5E-7Ga 6SCCM N2
GNSA-9566	2.30E + 15	52.4	
GNSA-9567			
GNSA-9568	3.80E + 14	328	
GNSA-9569	-3.60E + 13	45.9	
GNSA-9570	1.60E + 15	40.5	H2 only, no thermal source
GNSA-9571			

Appendix A: List of growth conditions for GaN samples

Sample Number	Carrier Concentration	Mobility	Notes
GNSA-9572	-2.50E + 18	4.9	
GNSA-9573	-1.40E + 18	118	
GNSA-9574			
GNSA-9575	-6.30E + 17	14.1	
GNSA-9611			Ga flux dropped for 1/2 hr (1036-988degree)
GNSA-9664			First hr @805 on dial, remainder @760
GNSA-9665			Tsub fluctuates during growth, N source leaks
GNSA-9666			Source leaks @10e-5 torr-l/sec
GNSA-9667			
GNSA-9668			
GNSA-9669			
GNSA-9670			

AN ATOMIC FORCE MICROSCOPY STUDY OF THE INITIAL NUCLEATION OF GaN ON SAPPHIRE

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ABSTRACT

Preliminary results of a study of GaN nucleation and growth by molecular beam epitaxy using a nitrogen rf plasma source are presented. Nucleation layers and 3000 Å thick layers were investigated by atomic force microscopy and x-ray diffraction. Growth under gallium-rich conditions both increased nucleation island size and promoted two-dimensional growth.

INTRODUCTION

The potential applications of blue and ultraviolet optoelectronic devices based on GaN have been recognized for many years [1]. Recent advances in epitaxial GaN growth by metal organic chemical vapor deposition (MOCVD) have lead to commercially available devices from both U.S. (CREE Research, Inc.) and foreign companies (Nichia Chemical Industries). Rapid progress in this direction is also being accomplished by molecular beam epitaxy (MBE) growth using active nitrogen species [2]. The most common substrate for epitaxial growth is sapphire (α - Al_2O_3) due to its availability, low cost and robust nature. GaN layers grown on sapphire, however, typically contain a high density of defects, mainly threading dislocations, due to a large lattice mismatch and thermal expansion coefficient mismatch between the epilayer and the substrate [3,4]. In contrast to growth in other semiconductor systems, these high dislocation densities ($>10^9 \text{ cm}^{-2}$) are reported to persist for up to 4 μm of growth [3,4]. While some devices are tolerant of such high densities [4], it is desirable to determine growth conditions for improved structural quality.

GaN typically nucleates and grows on sapphire by island formation. The use of low temperature buffer layers (450 - 600 °C) of AlN [5] or GaN [6,7] results in a dramatic improvement in layer morphology and electrical properties. Annealing prior to high temperature growth causes coalescence of the nucleation islands, resulting in low angle grain boundaries which create the observed dislocation arrays [3,4]. This subgrain structure is stable during growth under most conditions. Thus, subsequent crystal quality is strongly dependent on the nucleation layer.

The predominant growth mode is a further factor in dislocation reduction with increasing layer thickness. Two-dimensional growth results in the highest degree of structural perfection in epitaxial layer growth. Typical MBE and MOCVD growth conditions appear to promote three-dimensional growth [8]. This may lead to individual growth of the low angle grains, preventing dislocation recombination and annihilation. A recent study has reported MOCVD growth conditions resulting in two-dimensional step-flow growth [9], with a concomitant reduction in dislocation density to about $2 \times 10^8 \text{ cm}^{-2}$. This paper presents the initial results of a study to

determine conditions for MBE growth of GaN that result in buffer layers with increased grain size as well as determining the appropriate parameters to promote two-dimensional growth.

EXPERIMENT

The GaN layers for this study were grown at West Virginia University by MBE in a system similar to that described elsewhere [10]. Since we are interested in developing lower temperature growth of GaN by MBE, we have only investigated growth temperatures less than 700°C. A standard MBE source provided the Ga flux. A cryogenically-cooled rf plasma source (Oxford Applied Research CARS-25) operating at 500W was used to produce the active nitrogen flux. The layers were characterized using x-ray diffraction and atomic force microscopy (AFM) (Digital Instruments Nanoscope II).

RESULTS

Several studies [3,4,11] have indicated that the transition between buffer layer and "bulk" film structure occurs in the first 0.4 to 0.5 μm of growth. We grew a series of 3000 Å layers to determine both growth rates and growth modes in this transition region. We were also interested in the transition point between Ga-rich growth and conditions which produced Ga condensation as evidenced by the presence of Ga droplets. Figure 1 illustrates our growth rate for several conditions. Above 2.5×10^{-7} Torr Ga and 600°C, growth is limited by the amount of active nitrogen present as indicated both by the increase in growth rate with temperature at a fixed Ga flux, and by the relatively constant growth rate at a given temperature for increasing Ga flux. The decrease in growth rate between 2.5 and 5.0×10^{-7} Torr Ga at 660°C is apparently related to a change from a three-dimensional to a two-dimensional growth mode, as discussed later.

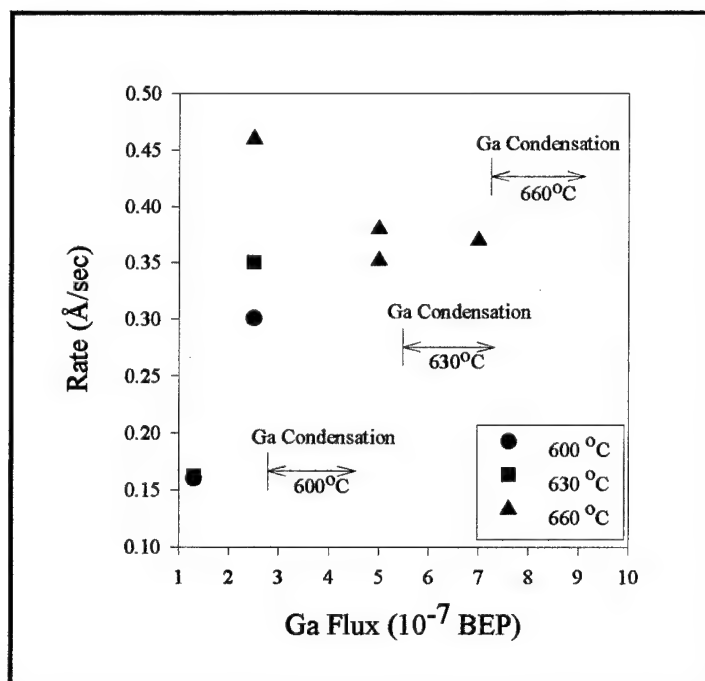


Figure 1. Growth rate of GaN for several temperatures and value of Ga flux.

Early in this study we grew layers using nucleation conditions reported by others [2,5,6,7,12]. We found that exposing the sapphire substrate to an active nitrogen flux resulted in a fine-grained ($<1000\text{\AA}$) island size. Growth of a low temperature nucleation layer at $450\text{ }^{\circ}\text{C}$ followed by an anneal at $660\text{ }^{\circ}\text{C}$ also resulted in a similar small grain size. Thus, we undertook a study of island size distributions in nucleation layers as a function of growth parameters. The nucleation layers were studied using AFM. Figure 2 is a micrograph of one such layer. Distributions were determined by taking ten AFM micrographs at points distributed across the sample surface. The islands were approximated as circular regions of various sizes by visual comparison with a template. A histogram of occurrence frequency vs. diameter was thus obtained. The distributions were adequately represented by poisson statistics, and the mean value was found by a least squares fit to the distribution.

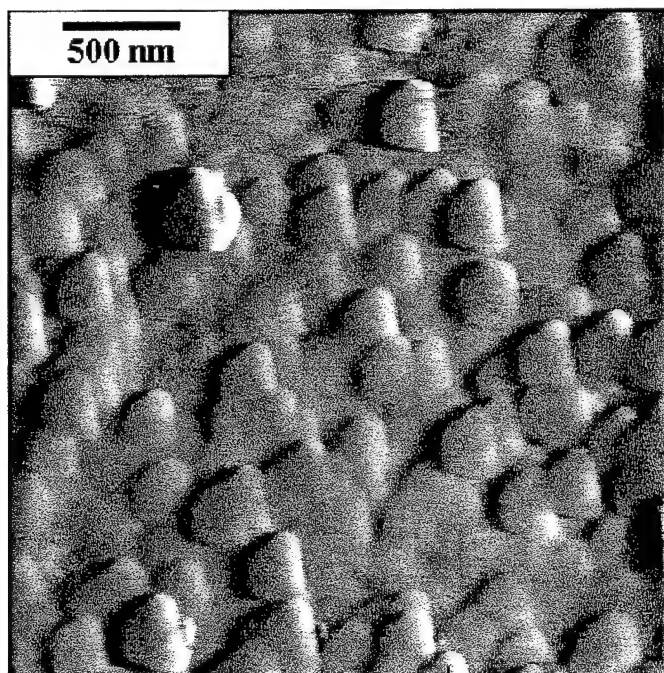


Figure 2. AFM micrograph of a GaN nucleation layer, 100\AA thick, grown at $660\text{ }^{\circ}\text{C}$ under 6 sccm nitrogen flow and 5.0×10^{-7} Torr BEP Ga. The average surface roughness of this layer was about 10 \AA . The mean island diameter was about 3000 \AA .

The nucleation island diameter was found to increase somewhat linearly with temperature over the range investigated. However, in order to obtain this increase, the nucleation had to occur under Ga-rich conditions. Indeed, at a given temperature, the largest island size always occurred near the boundary for Ga condensation. Our results are summarized in Figure 3. For a fixed Ga flux and nitrogen flow rate, we observed a temperature corresponding to a maximum island size. Above this temperature, the nucleation islands became smaller. At a fixed temperature, island size could be increased by increasing the Ga flux, up to the occurrence of Ga condensation. In contrast, increasing the nitrogen flow rate resulted in smaller island size.

Further information was gained by examining layers grown by extending the nucleation growth to 3000 \AA thick. Figure 4(a) is the AFM micrograph of such a layer grown at $660\text{ }^{\circ}\text{C}$. The morphology appeared to consist of well-defined three-dimensional microcrystallites, with an average surface roughness of about 150 nm . These growth parameters appear to promote three-dimensional growth without early coalescence of the islands. Also, up to the 3000 \AA thickness, there was no apparent change in average microcrystallite dimension from the original nucleation island size. X-ray diffraction measurements were consistent with single-crystal, hexagonal GaN for all layers measured. However, the full width at half maximum (FWHM) was fairly large, about 400 arc minutes for the sample depicted in Figure 4(a). Doubling the Ga-flux brought the growth to near-Ga-condensation conditions and gave an increase in island size, as seen in Figure

4(b). More importantly, however, is evidence of two-dimensional growth which was observed for all layers grown above 660 °C for Ga-rich conditions. Coalescence of the islands is occurring and the overall roughness between grains is reduced to about 20 nm. The tops of the islands are fairly flat with well defined sub-nanometer steps corresponding to a few monolayers. We believe this indicates we are near the conditions needed for two-dimensional step-flow growth. The x-ray diffraction FWHM was reduced to about 120 arc minutes for this layer.

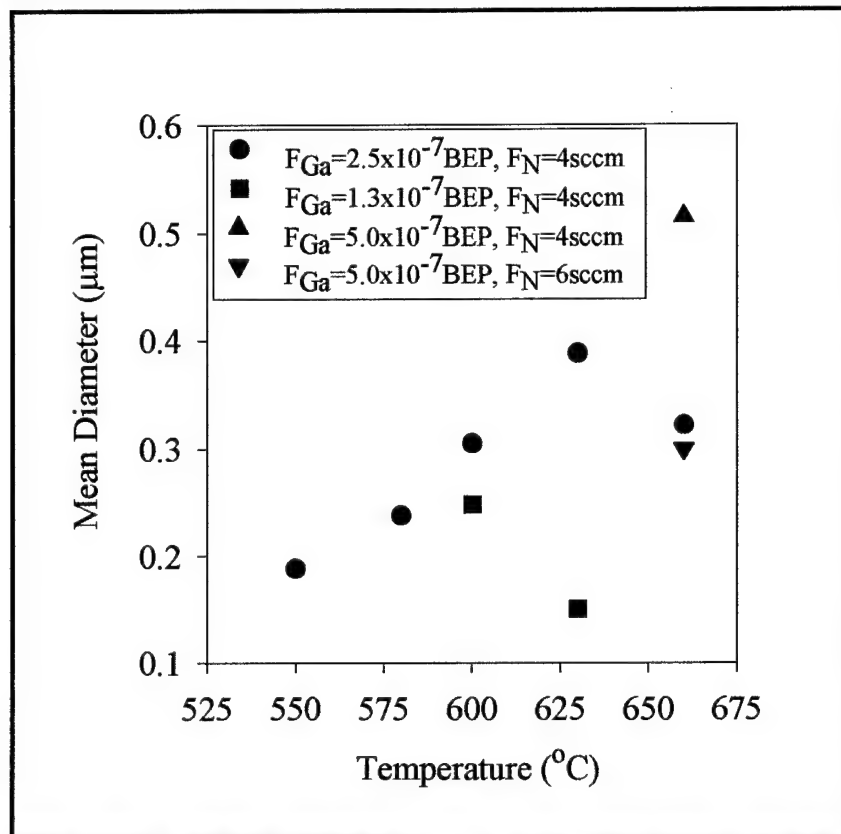


Figure 3 . GaN nucleation island diameter as a function of growth conditions.

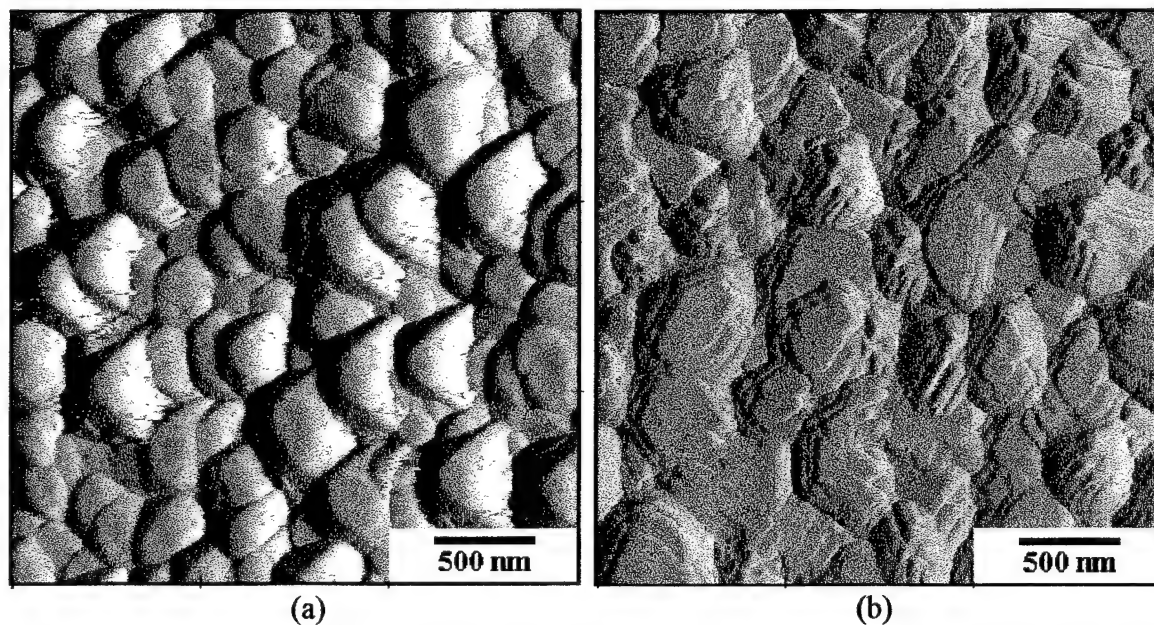


Figure 4 . AFM micrographs of GaN layers, 3000 Å thick, grown at 660°C under 4 sccm nitrogen flow and (a) 2.5×10^{-7} Torr BEP Ga (b) 5.0×10^{-7} Torr BEP Ga flux.

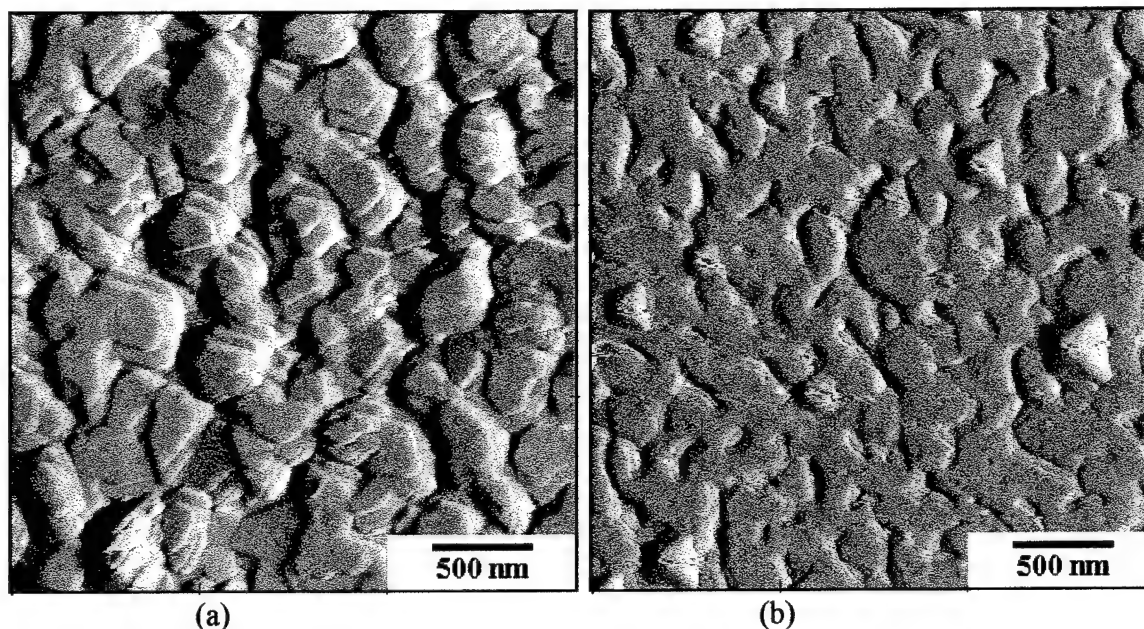


Figure 5 . AFM micrographs of GaN layers, 3000 Å thick, grown at 670°C under 6 sccm nitrogen flow and 5.0×10^{-7} Torr BEP Ga flux: (a) without a buffer layer and (b) with a 100 Å thick buffer layer annealed at 670°C for 20 minutes prior to layer growth.

Only layers grown above 660 °C under Ga-rich conditions, where increasing the Ga-flux does not increase the growth rate as shown in Figure 1, exhibit characteristics of two-dimensional growth. All others exhibited a well defined, three-dimensional microcrystallite structure. Figure 5(a) is an AFM micrograph of a layer grown at 670 °C under increased nitrogen flow to move further away from the Ga-condensation point while maintaining Ga-rich conditions. Coalescence is not as evident as for the previous sample, with an average surface roughness of about 30 nm. However, the islands are again flat-topped and steps corresponding to monolayer growth are observed. Figure 5(b) is a 3000 Å layer grown under the same conditions except that growth was interrupted after the first 100 Å of growth. This nucleation layer was then annealed at 670 °C for 20 minutes under nitrogen flux, and growth was resumed. The resulting layer exhibited almost complete coalescence, with an average surface roughness of about 1.5 nm. Sub-nanometer terraces were present on the top of the "flat" regions, again indicating a predominantly two-dimensional growth mode. The x-ray diffraction FWHM obtained for this layer was 51 arc minutes. Further optimization should result in complete coalescence and two-dimensional growth.

CONCLUSIONS

Apparently Ga is the more mobile species under our growth conditions since we obtained larger island sizes and two-dimensional growth only under Ga-rich conditions. Lowering the Ga flux lead to three-dimensional growth with a nucleation layer consisting of smaller island sizes. The nitrogen may only incorporate at available sites near where it initially adsorbs, with little lateral motion before desorbing. The best conditions for growing the nucleation layer appear to be as Ga-rich as possible, near the limit for Ga condensation.

In this study, we have determined conditions for MBE growth of GaN that give large nucleation island size, and promote two-dimensional growth of layers. However, the results presented here are preliminary, as we have not yet performed detailed optical and electrical characterizations. We plan to continue this study by growing thicker layers on annealed buffer layers with the larger island size for further characterization of material properties.

ACKNOWLEDGMENTS

We want to acknowledge the invaluable assistance of T. R. Charlton and D. Buczkowski in the counting of nucleation islands, and E. Hopkins for x-ray diffraction measurements. This work was supported by DoD/ONR Grant N00014-94-1-1149, monitored by M. Yoder.

REFERENCES

1. See, for example, J. Pankove, Mater. Res. Soc. Symp. Proc. **97**, 409 (1987).
2. T. D. Moustakas, T. Lei, and R. J. Molnar, Physica B **185**, 36 (1993).
3. W. Qian, M. Skowronski, M. DeGraef, K. Doverspike, L. B. Rowland, and D. K. Gaskill, Appl. Phys. Lett. **66**, 1252 (1995).
4. S. D. Lester, F. A. Ponce, M. G. Craford, and D. A. Steigerwald, Appl. Phys. Lett. **66**, 1249 (1995).
5. H. Amano, N. Sawaki, I. Asaki, and Y. Toyoda, Appl. Phys. Lett. **48**, 353 (1986).
6. S. Nakamura, Jpn. J. Appl. Phys. **30**, 1705 (1991).
7. J. N. Kuznia, M. Asif Khan, D. T. Olson, R. Kaplan, and J. Freitas, J. Appl. Phys. **73**, 4700 (1993).
8. Z. Sitar, L. L. Smith, and R. F. Davis, J. Cryst. Growth **141**, 11 (1994).
9. D. Kapolnek, X. H. Wu, B. Heying, S. Keller, B. P. Keller, U. K. Mishra, S. P. DenBaars, and J. S. Speck, Appl. Phys. Lett. **67**, 1541 (1995).
10. T. H. Myers, R. W. Yanka, K. A. Harris, A. R. Reisinger, J. Han, S. Hwang, Z. Yang, N. C. Giles, J. W. Cook, Jr., J. F. Schetzina, R. W. Green and S. McDevitt, J. Vac. Sci. Technol. **A7**, 300 (1989).
11. T. W. Weeks, Jr., M. D. Bremser, K. S. Ailey, E. Carlson, W. G. Perry, and R. F. Davis, Appl. Phys. Lett. **67**, 401 (1995).
12. M. E. Lin, B. N. Sverdlov, and H. Morkoç, J. Appl. Phys. **74**, 5038 (1993).

The effect of atomic hydrogen on the growth of gallium nitride by molecular beam epitaxy

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GaN was grown by molecular beam epitaxy using an rf plasma source. Growth under gallium-rich conditions at 730 °C was required to produce high quality layers as indicated by photoluminescence, Hall effect, atomic force microscopy, and x-ray diffraction measurements. Atomic hydrogen has a significant effect for Ga-rich growth, increasing growth rates by as much as a factor of 2. © 1996 American Institute of Physics. [S0003-6951(96)04344-6]

The potential of blue and ultraviolet optoelectronic devices has been recognized for many years.¹ Recent advances in epitaxial GaN growth by metalorganic chemical vapor deposition (MOCVD) have lead to the report of the first demonstration of a blue laser based on GaN.² Rapid progress in this direction is also being accomplished by molecular beam epitaxy (MBE) growth using active nitrogen species.^{3,4} One aspect of the MBE growth of GaN commonly cited as a disadvantage is the low growth rate. A recent study⁵ has reported the use of ammonia for the MBE growth of GaN, leading to growth rates comparable to that found in MOCVD growth.

Most studies have either ignored the effect of hydrogen on the growth, or have only considered its effects in compensating *p*-type dopants such as Mg.⁶ For example, the intent of using ammonia for MBE growth in the previously mentioned study was to obtain a larger flux of active nitrogen through catalytic decomposition on the GaN surface⁷ with hydrogen considered an undesirable by-product. Hydrogen may, however, be altering the growth kinetics as well. In this letter, we present evidence that atomic hydrogen can have a significant effect on the growth kinetics of GaN when the growth is limited by the amount of active nitrogen present.

The GaN layers for this study were grown at West Virginia University by MBE. A standard MBE source (EPI-40M) provided the Ga flux. A cryogenically cooled rf plasma source (Oxford Applied Research CARS-25) operating at either 500 or 600 W was used to produce the active nitrogen flux. All layers reported here were grown with a nitrogen flow rate of 6 sccm, resulting in a system background pressure of 6×10^{-5} Torr. Our growth rates were limited to less than 1 Å/s by the maximum flux of active nitrogen available from our source configuration. The distance between our nitrogen source and the growing layer was about 30 cm, nearly double that found in most MBE systems.

Atomic hydrogen was produced using a thermal cracker (EPI). Typically, 1×10^{-6} Torr beam equivalent pressure (BEP) of hydrogen was passed through the thermal source operating at 9.5 A, giving a dissociation efficiency of about 5% for this operating condition. Photoluminescence (PL)

measurements were performed using the 325 nm He-Cd laser line focused to a power density of 5 W/cm², a grating monochromator, and standard detection electronics. Hall, x-ray diffraction, and atomic force microscopy (AFM) (Digital Instruments Nanoscope II) measurements were also made.

Determination of substrate temperature and growth rate are important in this study. A calibrated spring-loaded type *K* thermocouple was in intimate contact with the back of the molybdenum sample block with the substrate mounted with indium. This calibration was routinely checked by using the melting points of various metals, oxide desorption from GaAs, and the use of an optical pyrometer for the higher temperatures. Temperature determination was reproducible to ± 5 °C.

Growth rate was determined using total sample thickness and the growth time. The 1 cm² sapphire substrates were held by a retaining mask which provided a step allowing measurement with a stylus surface profilometer. In addition, the thickness at the sample's center was determined from interference fringes in optical transmittance measured using a Cary-14 spectrophotometer. The edge and center thicknesses agreed to within 10%.

All samples were grown on *c*-plane sapphire substrates (Union Carbide Crystal Products). Prior to growth, the substrates were degreased and etched in a phosphoric/sulfuric (1:3) acid mixture heated to 80 °C. Based on our earlier study,⁸ buffer layers were grown by heating the substrate to 730 °C under an atomic hydrogen flux for 20 min and then cooling to 630 °C for the growth of a 200-Å-thick GaN buffer layer under a Ga flux of 5.0×10^{-7} Torr (BEP) with a 6 sccm nitrogen flow at 500 W. Buffer layer growth was initiated by simultaneous exposure to the Ga and N flux. This nucleation layer was then annealed at 730 °C for 20 min under nitrogen flux, cooled to the growth temperature, and growth was resumed. As discussed later, these conditions represent buffer layer growth under highly Ga-rich conditions. However, after the 730 °C anneal, examination of the buffer layers by AFM indicated continuous films with no evidence of Ga condensation.

GaN layers were grown under a variety of conditions, with the resulting growth rates indicated in Fig. 1. In Fig. 1, the growth rate is plotted as a function of Ga beam equivalent pressure. For an active nitrogen flux in excess or equal to the Ga flux, one would expect a linear increasing growth

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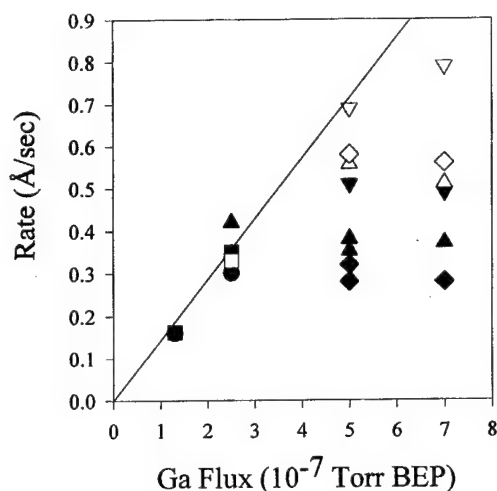


FIG. 1. Growth rate of GaN as a function of Ga flux. Substrate temperatures and rf power are: (●) 600 °C and 500 W; (■) 630 °C and 500 W; (▲) 660 °C and 500 W; (▼) 660 °C and 600 W; (◆) 730 °C and 600 W; (□) 630 °C and 500 W with H; (△) 660 °C and 500 W with H; (▽) 660 °C and 600 W with H; and (◇) 730 °C and 600 W with H.

rate with increasing Ga flux. When the surface ratio of Ga to active nitrogen is increased beyond unity, then either Ga condensation will occur if the desorption rate for excess Ga is low or the growth rate will become fairly constant with increasing Ga flux if the excess Ga desorption rate is high. A linear increase in growth rate was observed in going between 1.3 and 2.5×10^{-7} Torr BEP Ga. At growth temperatures of 600 (●) and 630 °C (■) with 500 W rf power, Ga-droplet condensation occurred for growths with a Ga flux above 2.5×10^{-7} Torr BEP Ga. In contrast, Ga condensation did not occur until beyond 7.0×10^{-7} Torr BEP upon raising the substrate temperature to 660 °C (▲), also at 500 W. The growth rate remained fairly constant for a Ga flux between 2.5 and 7.0×10^{-7} Torr BEP, indicating that the growth was limited by the amount of active nitrogen available.

Increasing the power into the nitrogen rf source increases the amount of active nitrogen.⁹ The concomitant increase in growth rate observed in going from 500 (▲) to 600 W (▼) power at 660 °C thus supports the assertion that the growth rate is limited by the amount of active nitrogen present. Our study is also in agreement with others,¹⁰ that temperatures above 700 °C are necessary to obtain high quality material, as detailed below. Unfortunately, the GaN growth rate was observed to decrease significantly in going from 660 °C (▼) to 730 °C (◆) at 600 W rf power, from about 0.5 to about 0.3 Å/s. The increase in temperature has apparently either reduced the residence time of unreacted N on the surface or increased the Ga desorption rate, thereby reducing the growth rate. Prior studies of Ga desorption for GaN growth using ammonia¹¹ indicate that the Ga desorption rate increases rapidly above 700 °C.

The surface morphology was distinctly different for layers grown under Ga-rich conditions as compared to layers grown closer to equal Ga and active nitrogen flux. Nitrogen sufficient conditions gave a highly textured, three-dimensional surface, as shown by the micrograph in Fig. 2(a). Such morphology is similar to that reported earlier¹² for the growth of GaN by MBE using an identical rf nitrogen

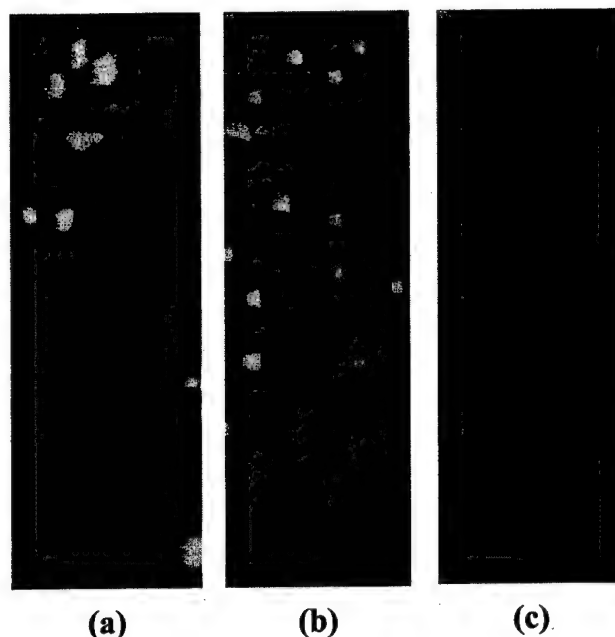


FIG. 2. AFM micrographs ($5 \mu\text{m} \times 15 \mu\text{m}$) illustrating the typical surface morphologies for: (a) nearly equal gallium and active nitrogen flux, (b) intermediate gallium flux, and (c) gallium-rich conditions. The height scale is 135, 200, and 35 nm, respectively, for the micrographs.

source. Increasing the Ga flux resulted in isolated structures interspersed on flat regions of the surface, as can be seen in Fig. 2(b). The measured rms. surface roughness between the structures ranged between 0.6 and 1.5 nm. As we reported earlier,⁸ Ga-rich growth close to, but below the Ga condensation point resulted in smooth, featureless surfaces indicative of two-dimensional growth with an rms surface roughness of 1.5 nm over a $15 \mu\text{m}$ span, similar to that reported for high quality MOCVD growth.¹³ This can be seen in Fig. 2(c). Subnanometer tall terraces representing atomic steps were present on the top of the "flat" regions of the sample.

This change in surface morphology is very similar to that reported earlier¹⁴ for growth of GaN using an electron cyclotron resonance microwave (ECR) plasma source. In the ECR study, the amount of active nitrogen was changed by controlling the plasma density. Lower power led to smooth surfaces, while high powers resulted in textured surfaces. It is known that, in addition to increasing the active nitrogen flux, high-power operation of an ECR source can lead to a significant amount of high-energy ions in the nitrogen flux¹⁵ which could influence the growth morphology. In our study, the nitrogen source parameters were unchanged for the samples shown in Fig. 2 (600 W, 6 sccm) while the Ga flux was altered. Thus, the agreement between the two studies suggests that the change from smooth to textured growth is a universal phenomenon related to the ratio of active nitrogen to gallium at the growth front.

Introduction of atomic hydrogen resulted in a dramatic increase in the growth rate of the GaN under Ga-rich conditions as also summarized in Fig. 1. The samples whose growth rates are indicated by the open symbols in Fig. 1 were grown under a total (atomic and molecular) hydrogen flux of 1×10^{-6} Torr BEP. The other growth parameters remained the same. Note that the growth rate was essentially

doubled for each case of Ga-rich growth. In contrast, a sample grown under nitrogen-sufficient conditions at 630 °C did not exhibit an enhanced growth rate. The increase in growth rate for the Ga-rich cases did not appear to be very sensitive to the overall hydrogen flux. Changing the hydrogen flux from 0.5 to 2.0×10^{-6} Torr BEP gave the same value for the increase in the growth rate. To see if the increased growth rate originated with molecular hydrogen, samples were grown under hydrogen flux with the cracker turned off. The resulting samples exhibited identical growth rates to the GaN grown without hydrogen, indicating that molecular hydrogen is not significantly affecting the growth kinetics. Also, to see if the atomic hydrogen was possibly forming active species with molecular nitrogen, an attempt was made to grow under an atomic hydrogen flux with the rf power turned off on the nitrogen source. The resulting GaN growth rate, if nonzero, was too small to be detected.

The increase in growth rate for Ga-rich conditions is apparently related to the presence of atomic hydrogen. We propose that the atomic hydrogen becomes loosely bonded to the growing GaN surface. Nitrogen atoms adsorbed on the surface are then attracted by this hydrogen layer, resulting in an increased nitrogen residence time. The longer residence time increases the probability that a Ga atom will diffuse to within an interaction distance of the nitrogen, and thus enhance the growth rate of GaN. Thus, the atomic hydrogen could be increasing the effective active nitrogen concentration. In addition, the surface morphology for samples grown under atomic hydrogen more nearly resembled that shown as Fig. 2(a). This, along with the increased growth rate, is consistent with the growth kinetics towards a more nitrogen-sufficient case.

PL measurements were made at both room and liquid helium temperatures on the samples. At room temperature, samples investigated exhibited band-edge emission at about 365 nm (3.39 eV). All samples grown at or below 660 °C also exhibited significant luminescence centered at about 560 nm (2.2 eV). Layers grown at 730 °C did not exhibit detectable yellow luminescence. We take this as direct evidence that the higher growth temperatures are necessary to produce high quality material. In low-temperature PL measurements, the layers grown at 730 °C exhibited primarily near-band-edge luminescence related to excitonic transitions, again indicating high quality layers. The PL obtained from a GaN layer grown at 730 °C under an atomic hydrogen flux is shown in Fig. 3. There is a single bound-exciton peak at about 3.47 eV and no evidence of PL related to deep levels, indicating that growth under hydrogen does not degrade layer quality.

The best GaN samples grown without atomic hydrogen were grown under Ga-rich conditions at 730 °C and exhibited x-ray diffraction rocking curve full widths at half maximum (FWHM) between 4 and 5 arcmin. Hall measurements made on these layers indicated *n*-type carrier concentrations as low as $4 \times 10^{17} \text{ cm}^{-3}$ with room-temperature mobilities as high as $120 \text{ cm}^2/\text{V s}$. The x-ray FWHM of the samples grown with atomic hydrogen were indistinguishable from those grown without atomic hydrogen under the same conditions. However, the background *n*-type carrier concentration was about a factor of 2 larger, at the low 10^{18} cm^{-3}

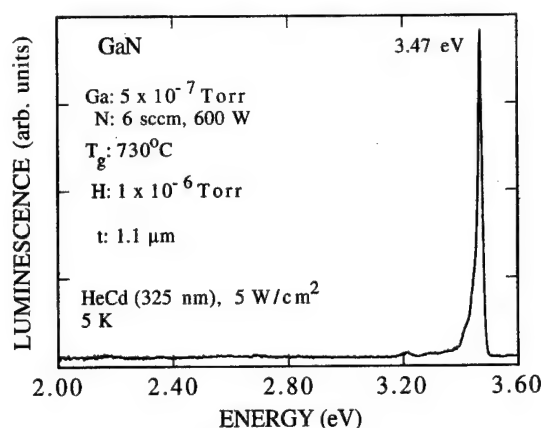


FIG. 3. Photoluminescence of GaN.

level also with a mobility of about $120 \text{ cm}^2/\text{V s}$. Although hydrogen may introduce donor levels in GaN,¹⁶ we have no direct evidence that this is the case here. As shown in Fig. 3, we did not observe the 3.35 eV PL feature which has been associated with the hydrogen-related donor.¹⁶ It may be that the higher carrier concentration is related to the three-dimensional growth indicated by the textured surface.

In conclusion, we have demonstrated that the presence of atomic hydrogen can have a significant effect on the growth rate of GaN under Ga-rich conditions. PL and Hall measurements indicate that layer quality is not degraded by growth under atomic hydrogen. The increased growth rate and change to a textured surface morphology are suggestive that the atomic hydrogen increases the effective surface concentration of nitrogen. In addition, we have presented evidence that Ga-rich growth promotes a smoother growth surface.

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¹See, for example, J. Pankove, *Mater. Res. Soc. Symp. Proc.* **97**, 409 (1987).

²S. Nakamura, M. Senoh, S. Nagahama, N. Iwasa, T. Yamada, T. Matsushita, H. Kiyoku, and Y. Sugimoto, *Jpn. J. Appl. Phys.* **35**, L74 (1996).

³T. D. Moustakas, T. Lei, and R. J. Molnar, *Physica B* **185**, 36 (1993).

⁴M. Smith, G. D. Chen, J. Z. Li, J. Y. Lin, H. X. Jiang, A. Salvador, W. K. Kim, O. Aktas, A. Botchkarev, and H. Morkoc, *Appl. Phys. Lett.* **67**, 3387 (1995).

⁵Z. Yang, L. K. Li, and W. I. Wang, *Appl. Phys. Lett.* **67**, 1686 (1995).

⁶M. S. Brandt, J. W. Ager III, W. Gotz, N. M. Johnson, J. S. Harris, Jr., R. J. Molnar, and T. D. Moustakas, *Phys. Rev. B* **49**, 14758 (1994).

⁷S. S. Liu and D. A. Stevenson, *J. Electrochem. Soc.* **125**, 1161 (1978).

⁸M. Richards-Babb, S. L. Buczkowski, Zhonghai Yu, and T. H. Myers, *Mater. Res. Soc. Symp. Proc.* **395**, 237 (1996).

⁹R. P. Vaudo, J. W. Cook, Jr., and J. F. Schetzina, *J. Cryst. Growth* **138**, 430 (1994).

¹⁰T. D. Moustakas, *Mater. Res. Soc. Symp. Proc.* **395**, 111 (1996).

¹¹C. R. Jones, T. Lei, R. Kaspi, and K. R. Evans, *Proceedings of the 1995 Fall MRS Meeting, Symposium AAA* (unpublished).

¹²C. Trager-Cowan, K. P. O'Donnell, S. E. Hooper, and C. T. Foxon, *Appl. Phys. Lett.* **68**, 355 (1996).

¹³D. Kapolnek, X. H. Wu, B. Heying, S. Keller, B. P. Keller, U. K. Mishra, S. P. DenBaars, and J. S. Speck, *Appl. Phys. Lett.* **67**, 1541 (1995).

¹⁴T. D. Moustakas and R. J. Molnar, *Mater. Res. Soc. Symp. Proc.* **281**, 753 (1993).

¹⁵R. J. Molnar and T. D. Moustakas, *J. Appl. Phys.* **76**, 4587 (1994).

¹⁶M. S. Brandt, N. M. Johnson, R. J. Molnar, R. Singh, and T. D. Moustakas, *Appl. Phys. Lett.* **64**, 2264 (1996).

THE EFFECT OF HYDROGEN ON THE MOLECULAR BEAM EPITAXY GROWTH OF GaN ON SAPPHIRE UNDER Ga-RICH CONDITIONS

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ABSTRACT

Nucleation and growth of GaN under Ga-rich conditions by molecular beam epitaxy using a nitrogen rf plasma source is shown to result in both a smoother GaN surface and a reduced inversion domain content. In addition, preliminary results of the dramatic effect of atomic hydrogen on growth kinetics for Ga-rich growth are presented.

INTRODUCTION

GaN typically nucleates and grows on sapphire by island formation. The use of low temperature buffer layers (450 - 600 °C) of AlN [1] or GaN [2,3] results in a dramatic improvement in layer morphology and electrical properties. Annealing prior to high temperature growth causes coalescence of the nucleation islands, resulting in low angle grain boundaries which create the observed dislocation arrays [4,5]. It appears that this subgrain structure is stable during growth under most conditions. Thus, subsequent crystal quality is strongly dependent on the nucleation layer. The predominant growth mode is a further factor in dislocation reduction with increasing layer thickness. Two dimensional growth results in the highest degree of structural perfection in epitaxial layer growth. Typical molecular beam epitaxy (MBE) and metal-organic chemical vapor deposition (MOCVD) growth conditions appear to promote three dimensional growth of GaN. [6,7] This may lead to individual growth of the low angle grains, preventing dislocation recombination and annihilation. A recent study has reported MOCVD growth conditions resulting in two-dimensional step-flow growth [8], with a concomitant reduction in dislocation density to about $2 \times 10^8 \text{ cm}^{-2}$. It is desirable to achieve conditions for MBE growth to allow such a growth mode to occur.

One of the obvious differences between MOCVD and MBE growth of GaN is the presence of hydrogen, primarily as a component of the molecule supplying the nitrogen. Most studies have either ignored the effect of hydrogen, or have only considered its effects in compensating p-type dopants such as Mg. For example, the use of ammonia in MBE growth is primarily to obtain a larger flux of active nitrogen through catalytic decomposition on the GaN surface, again with the hydrogen considered only as a reaction by-product. In this paper, we present evidence that: nucleation under Ga-rich conditions results in reduced dislocation and inversion domain content; continued growth under Ga-rich conditions promotes a smoother, two-dimensional growth front; and hydrogen can have a significant effect on the growth kinetics of GaN when the growth is limited by the amount of active nitrogen present.

EXPERIMENT

The GaN layers for this study were grown at West Virginia University by MBE in a system described elsewhere. [9] A standard MBE source provided the Ga flux. A cryogenically-cooled rf plasma source (Oxford Applied Research CARS-25) operating at 600W was used to produce active nitrogen flux. Atomic hydrogen was produced using a commercial thermal

cracker (EPI-AHS). Our source-to-substrate distance was large, about 14 inches, leading to a lower total active-nitrogen flux than typically obtained from an rf plasma source. When scaled by the differences in source-to-substrate distance ($1/r^2$), the active nitrogen flux was comparable to that reported by others. The layers were characterized by Hall measurements, photoluminescence, x-ray diffraction, and atomic force microscopy (AFM) (Digital Instruments Nanoscope II). High resolution transmission electron microscopy was performed at Xerox Palo Alto Research Center using techniques detailed elsewhere [10]

RESULTS

All samples were grown on c-plane sapphire substrates (Union Carbide Crystal Products). Prior to growth, the substrates were degreased and etched in a phosphoric/sulfuric (1:3) acid mixture heated to 80°C. Based on our earlier study,[11] buffer layers were grown by heating the substrate to 730 °C under an atomic hydrogen flux for 20 minutes and then cooling to 630 °C for the growth of a 200 Å thick GaN buffer layer under a Ga flux of 5.0×10^{-7} Torr (BEP) with a 6 SCCM nitrogen flow at 600 W. This procedure results in nucleation island sizes around 0.3 to 0.5 μm, giving dislocations due to domain-wall coalescence of order 10^9 cm^{-2} . Buffer layer growth was initiated by simultaneous exposure to the Ga and N flux. The 20 nm nucleation layer was then annealed at 730 °C for 20 minutes under active nitrogen flux, cooled (or heated) to the growth temperature, and growth was resumed. Our conditions represented highly Ga-rich growth for the buffer layer. However, after the 730 °C anneal, examination of buffer layers by AFM indicated continuous films with no evidence of Ga condensation.

GaN layers were grown on the annealed buffer layers. In Fig. 1, the growth rate is plotted as a function of Ga beam equivalent pressure. For an active nitrogen flux in excess or equal to the Ga flux (nitrogen sufficient), one would expect a linear increase in growth rate with increasing Ga flux. This is represented by the solid line in the figure. When the surface ratio of Ga to active nitrogen is increased beyond unity, then either Ga condensation will occur if the desorption rate for excess Ga is low

or the growth rate will become fairly constant with increasing Ga flux if the excess Ga desorption rate is large enough to prevent condensation. The surface morphology was distinctly different for layers grown under Ga-rich conditions as compared to layers grown closer to equal Ga and active nitrogen flux. Nitrogen sufficient conditions gave a highly textured, three-dimensional surface, as shown by the micrograph in Fig. 2(a). Such morphology is similar to that reported earlier [6] for the growth of GaN by MBE using an identical rf nitrogen source. Ga-rich growth close to, but

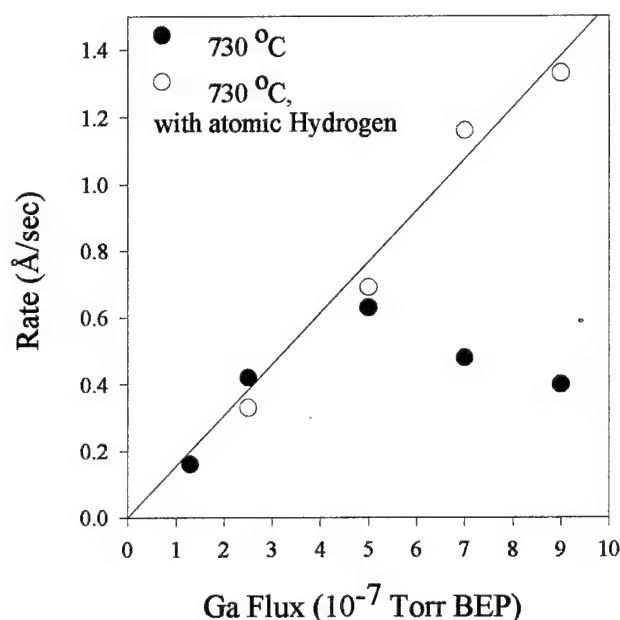


Figure 1. Growth rate v. Ga-flux.

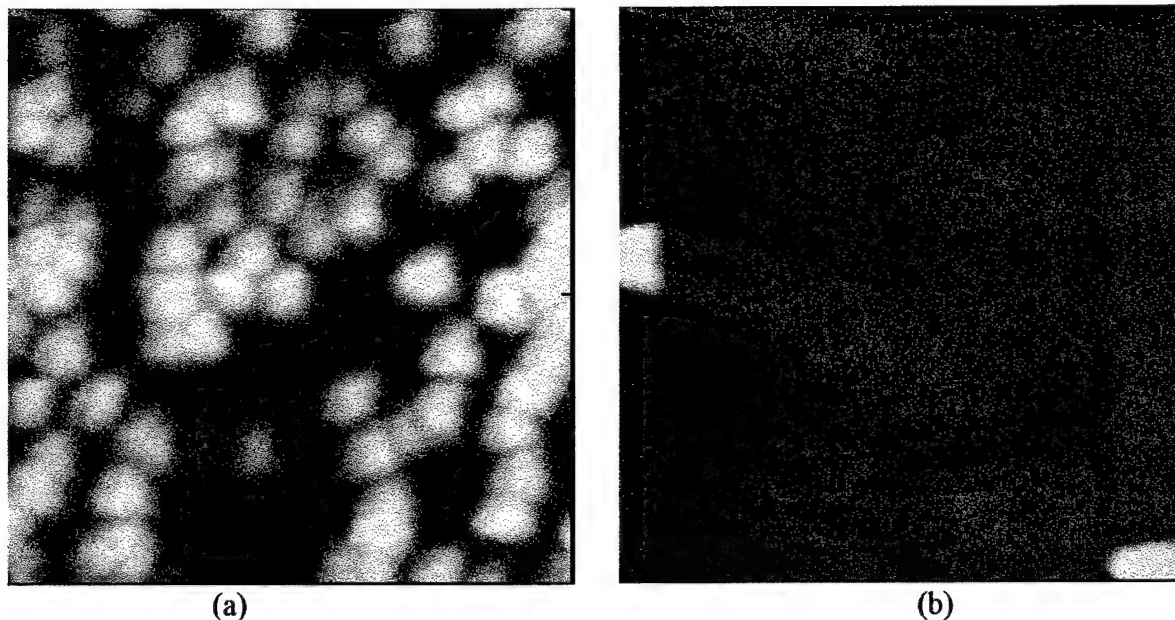


Figure 2. AFM micrographs of GaN grown under (a) nitrogen-sufficient and (b) Ga-rich conditions. The areas represented are $2 \text{ by } 2 \mu\text{m}^2$. The height scales are 135 nm in (a) and 50 nm in (b).

below the Ga condensation point resulted in much smoother surfaces indicative of two-dimensional growth with an rms. surface roughness of 1.5 nm over a $15 \mu\text{m}$ span. The smoother texture can be seen in the AFM micrograph in Fig. 2(b) for a 250 nm thick layer. Sub-nanometer tall terraces representing atomic steps were present on the top of the "flat" regions of the sample.

This change in surface morphology is very similar to that reported earlier [12] for growth of GaN using an electron cyclotron resonance microwave (ECR) plasma source. In the ECR study, the amount of active nitrogen was changed by controlling the plasma density. Lower power led to smooth surfaces, while high powers resulted in textured surfaces. It is known that, in addition to increasing the active nitrogen flux, high power operation of an ECR source can lead to a significant amount of high energy ions in the nitrogen flux which could influence the growth morphology. In our study, the nitrogen source parameters were unchanged for the samples shown in Fig. 2 while the Ga flux was altered. Thus, the agreement between the two studies suggests that the change from smooth to textured growth is a universal phenomenon related to the ratio of active nitrogen to gallium at the growth front.

The sample whose AFM micrograph is shown in Fig. 2(b) was also measured by high resolution TEM. As shown in Fig. 3, the microstructure is characterized by a high density of threading dislocations at the GaN-sapphire interface. These quickly annihilate resulting in a line defect density in the low 10^9 cm^{-2} after growth of only about 250 nm of GaN. By comparison, samples grown under nitrogen-sufficient conditions had line defect densities in the mid- 10^{10} cm^{-2} . This decrease can be attributed to the layer-by-layer growth mode indicated by the smooth surface morphology measured by AFM. Thicker layers should result in commensurably fewer dislocations. Perhaps of more significance, the sample exhibited a low concentration of inversion domain boundaries, where the GaN crystal structure is inverted along the c-axis due to nucleation of the opposite phase at the substrate. As reported previously, inversion domain boundaries are a significant source of defects in GaN grown by any technique. [10] Our direct nucleation of GaN on sapphire under Ga-rich conditions appears to suppress the formation of such inversion domains, resulting in significantly less than 10 % by volume. This is comparable to most

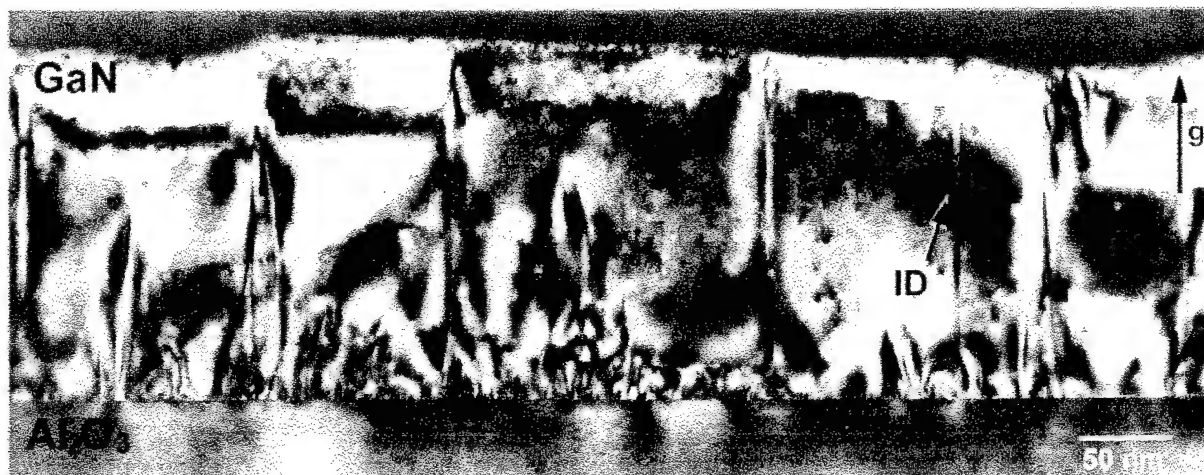


Figure 3. Dark Field TEM image taken with $g = (0002)$ near the $[11-20]$ zone axis of GaN. ID indicates the inversion domain in this micrograph. The remaining defects are nonedge-type dislocations.

MOCVD layers. In contrast, ECR-MBE layers grown on nitrided sapphire contain about an equal number of both orientations.[10]

Increasing the growth temperature should also increase the surface mobility of adatoms and promote better layer-by-layer growth, which may increase the rate of dislocation clean-up. Unfortunately, we [9] and others [13] have observed that the GaN growth rate decreases significantly at temperatures above about 730 °C. Further increase in temperature apparently either reduces the residence time of unreacted N on the surface or increases the Ga desorption rate, thereby reducing the growth rate. Prior studies of Ga desorption for GaN growth using ammonia [14] indicate that the Ga desorption rate increases rapidly above 730 °C. Thus, while this temperature can be increased somewhat by increasing the amount of active nitrogen, the regime between 750 and 800 °C may represent a practical limit to the growth temperature feasible by MBE using nitrogen plasma sources.

Introduction of atomic hydrogen resulted in a dramatic increase in the growth rate of the GaN under Ga-rich conditions, as also summarized in Fig. 1. The samples whose growth rates are indicated by the open symbols in Fig. 1 were grown under a total (atomic and molecular) hydrogen flux of 1 to 2×10^{-6} Torr BEP. The other growth parameters remained the same as for the corresponding filled-symbol case. Note that the growth rate was essentially brought up to the Ga-limit for each case of Ga-rich growth. In contrast, samples grown under nitrogen-sufficient conditions did not exhibit an enhanced growth rate. To see if the increased growth rate originated with molecular hydrogen, samples were grown under hydrogen flux with the cracker turned off. The resulting samples exhibited identical growth rates to the GaN grown without hydrogen, indicating that molecular hydrogen is not significantly affecting the growth kinetics. Also, to see if the atomic hydrogen was possibly forming active species with molecular nitrogen, an attempt was made to grow under an atomic hydrogen flux with the rf power turned off on the nitrogen source. The resulting GaN growth rate, if non-zero, was too small to be detected. Ammonia was not detected in an analysis of the background gases present during growth. This indicates that the enhanced growth rate is not due to the presence of ammonia formed due to gas-phase reactions.

The increase in growth rate for Ga-rich conditions is apparently related to the presence of atomic hydrogen at the surface of the growing layer. We propose that the atomic hydrogen becomes loosely bonded to the growing GaN surface. Nitrogen atoms adsorbed on the surface

are then attracted by this hydrogen layer, resulting in an increased nitrogen residence time. The longer residence time increases the probability that a Ga atom will diffuse to within an interaction distance of the nitrogen, and thus enhance the growth rate of GaN. Thus, the atomic hydrogen could be increasing the effective active nitrogen concentration. In addition, the surface morphology for samples grown under atomic hydrogen more nearly resembled that shown as Fig. 2(a). This, along with the increased growth rate, is consistent with shifting the growth kinetics towards a more nitrogen-sufficient case. Of practical importance, this shifting of the growth kinetics may allow much higher growth temperatures for MBE of GaN. We are continuing to investigate this phenomenon.

PL measurements were made at liquid helium temperatures on the samples. All samples grown at or below 660 °C exhibited significant luminescence centered at about 560 nm (2.2 eV). While the origin of this broad yellow luminescence band is still controversial, it is commonly attributed to deep states in the bandgap involving impurities or native defects. As reported earlier, [9] layers grown at 730 °C did not exhibit detectable yellow luminescence. We take this as direct evidence that the higher growth temperatures are necessary to produce high quality material. The PL obtained from a GaN layer grown at 660°C under an atomic hydrogen flux is shown in Fig. 4. There is a single bound-exciton peak at about 3.48 eV and a small amount of yellow luminescence. As with the undoped layers, raising the growth temperature above 730 °C eliminated the yellow PL for samples grown under hydrogen, [9] indicating that growth under hydrogen does not degrade layer quality.

The best GaN samples grown without atomic hydrogen were grown under Ga-rich conditions at 730 °C and exhibited x-ray diffraction rocking curve full widths at half maximum (FWHM) between 2 and 3 arc minutes for samples with thickness about 1 μm . Hall measurements indicated n-type carrier concentrations as low as $1 \times 10^{17} \text{ cm}^{-3}$ with room temperature mobilities of $120 \text{ cm}^2/\text{V}\cdot\text{s}$. The x-ray FWHM and Hall measurements of samples grown with atomic hydrogen were indistinguishable from those grown without atomic hydrogen. Although previously reported that hydrogen may introduce donor levels in GaN,[15] we have no direct evidence that this is the case here. As shown in Fig. 4 and in previously published spectra from samples grown at higher temperatures, we did not observe the 3.35 eV PL feature which has been associated with the hydrogen-related donor.[15] In addition, we did not see any significant difference in background doping between layers grown with and without hydrogen.

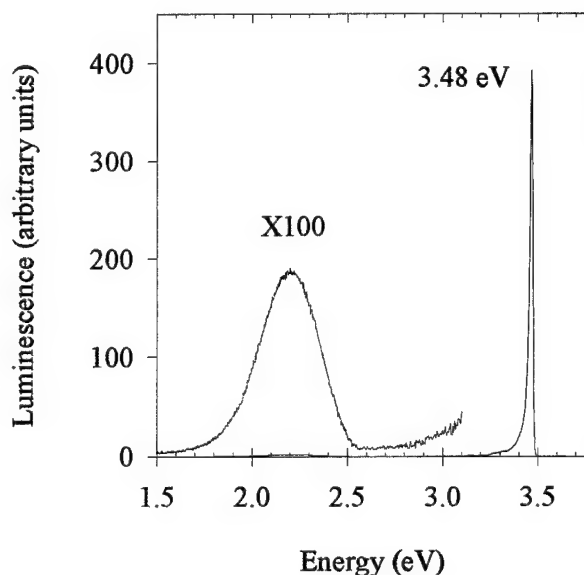


Figure 4. Photoluminescence of GaN grown at 660 °C under an atomic hydrogen flux.

CONCLUSIONS

In conclusion, we have demonstrated that the presence of atomic hydrogen can have a significant effect on the growth rate of GaN under Ga-rich conditions, resulting in an increase in growth rate limited by the total Ga flux. PL and Hall measurements indicate that layer quality is not degraded by growth under atomic hydrogen. The increased growth rate and change to a textured surface morphology are suggestive that the atomic hydrogen increases the effective surface concentration of nitrogen. In addition to the hydrogen-related growth rate enhancement, we have presented evidence that Ga-rich nucleation and growth results in a significant reduction of inversion domain boundaries, promotes dislocation reduction and promotes a smoother growth surface.

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REFERENCES

1. H. Amano, N. Sawaki, I. Asaki, and Y. Toyoda, *Appl. Phys. Lett.* **48**, 353 (1986).
2. S. Nakamura, *Jpn. J. Appl. Phys.* **30**, 1705 (1991).
3. J. N. Kuznia, M. Asif Khan, D. T. Olson, R. Kaplan, and J. Freitas, *J. Appl. Phys.* **73**, 4700 (1993).
4. W. Qian, M. Skowronski, M. DeGraef, K. Doverspike, L. B. Rowland, and D. K. Gaskill, *Appl. Phys. Lett.* **66**, 1252 (1995).
5. S. D. Lester, F. A. Ponce, M. G. Crafford, and D. A. Steigerwald, *Appl. Phys. Lett.* **66**, 1249 (1995).
6. C. Trager-Cowan, K.P. O'Donnell, S.E. Hooper and C.T. Foxon, *Appl. Phys. Lett.* **68**, 355 (1996).
7. Z. Sitar, L. L. Smith, and R. F. Davis, *J. Cryst. Growth* **141**, 11 (1994).
8. D. Kapolnek, X. H. Wu, B. Heying, S. Keller, B. P. Keller, U. K. Mishra, S. P. DenBaars, and J. S. Speck, *Appl. Phys. Lett.* **67**, 1541 (1995).
9. Zhonghai Yu, S.L. Buczkowski, N.C. Giles, T.H. Myers and M.R. Richards-Babb, *Appl. Phys. Lett.* **69**, 2731 (1996).
10. L.T. Romano, J.E. Northrup and M.A. O'Keefe, *Appl. Phys. Lett.* **69**, 2394 (1996).
11. M. Richards-Babb, S.L. Buczkowski, Zhonghai Yu, And T.H. Myers, *Mater. Res. Soc. Symp. Proc.* **395**, 237 (1996).
12. T.D. Moustakas and R.J. Molnar, *Mat. Res. Symp. Proc.* **281**, 753 (1993).
13. See, for example, S. Guha, N.A. Bojarczuk and D.W. Kisher, *Appl. Phys. Lett.* **69**, 2879 (1996).
14. C.R. Jones, T. Lei, R. Kaspi and K.R. Evans, *Mater. Res. Soc. Symp. Proc.* **395**, 141 (1996).
15. M.S. Brandt, N.M. Johnson, R.J. Molnar, R. Singh and T.D. Moustakas, *Appl. Phys. Lett.* **64**, 2264 (1996).

The influence of inversion domains on surface morphology in GaN grown by molecular beam epitaxy

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Growth of GaN by rf-plasma molecular beam leads to different surface morphologies for nitrogen-rich growth versus gallium-rich growth. Nitrogen-rich growth produces a significant density of pyramidal hillocks while gallium-rich growth results in flat surfaces. Differences in surface morphology were directly linked to the presence of inversion domains which originated in the nucleation layer. Nitrogen-rich growth and growth under atomic hydrogen enhanced the growth rate of inversion domains with respect to the surrounding matrix, while growth under Ga-rich conditions resulted in a more nearly equal growth rate. © 1997 American Institute of Physics. [S0003-6951(97)00350-1]

Heteroepitaxial GaN layers grown by any technique contain a high dislocation density as well as other classes of defects such as nanopipes, voids and inversion domain boundaries (IDBs). Inversion domains (IDs), consisting of regions of GaN with the opposite polarity to the primary matrix, have been observed to some extent in GaN grown by all techniques.¹⁻⁵ However, there is some indication that IDBs are both more common and more stable during growth of GaN by molecular beam epitaxy (MBE).^{5,6} In this letter, we present the results of a study of IDBs in GaN grown by rf-plasma-assisted MBE detailing how different growth conditions affect the role of IDBs in determining surface morphology.

The presence of IDs have been previously reported for GaN grown by MBE.^{5,6} Samples grown by electron cyclotron resonance (ECR) MBE nucleated under Ga-rich conditions exhibited IDs up to 50% by volume,⁵ which is significantly larger than that observed in GaN grown by metal-organic chemical vapor deposition (MOCVD).¹⁻³ While films nucleated and grown under nitrogen-rich (N-rich) conditions by ECR-MBE were found to be free of IDBs, significant regions of zincblende material near the interface were observed.⁵ Also, nucleation and growth of GaN under Ga-rich conditions has resulted in improved structural, electrical and optical properties with smoother surface morphologies when compared to N-rich growth.⁶⁻⁸

Atomic force microscopy (AFM) and transmission electron microscopy (TEM) were used to study surface morphology and its relation to microstructure in a series of GaN layers grown by MBE on (0001) sapphire. Detailed descriptions of the growth procedure and other layer characterizations are given elsewhere.^{6,9,10} In brief, the buffer layers for all samples were grown by heating the substrate to 730 °C under an atomic hydrogen flux for 20 min and then cooling to 630 °C for the growth of a 200-Å-thick GaN buffer layer.⁶ GaN layers were then grown at a temperature of 730 °C under various N-rich and Ga-rich conditions, both with and

without an atomic H flux. A standard Ga source and an Oxford CARS25 rf-plasma nitrogen source were used for the growths. Cross-section TEM (XTEM) studies were performed with a JEOL 3010 microscope operated at 300 kV on samples that were prepared by polishing and then ion milling to electron transparency. TEM images were taken under various diffraction conditions including multiple dark-field imaging with $g = \pm(0002)$ along either the $\langle 11\bar{2}0 \rangle$ or $\langle 10\bar{1}0 \rangle$ axis in order to reveal inversion domains.¹ AFM was performed in air using a Digital Instruments Nanoscope II.

As detailed by Yu *et al.*,⁶ AFM measurements indicate smooth surfaces for layers grown under Ga-rich conditions without atomic H, with rms roughness values < 1 nm. In contrast, samples grown either under N-rich conditions or under an atomic hydrogen flux have a rough surface consisting of pyramidal hillocks. The hillocks have a triangular cross section that is $\sim 0.5 \mu\text{m}$ on a side for $1\text{-}\mu\text{m}$ -thick samples, and between 150–250 nm in height. Growth near the N-rich/Ga-rich transition (without atomic H) results in smaller size pyramids. The surfaces between the hillocks have been measured by AFM and x-ray reflectivity to have a rms roughness of 1–2 nm.^{6,10}

Figure 1(a) shows a XTEM image of a $1.0\text{-}\mu\text{m}$ -thick sample grown under an atomic hydrogen flux with a surface that contains a high density of pyramid-shaped hillocks similar to that observed for all samples grown under nitrogen-rich conditions, or with atomic hydrogen. The hillocks are ~ 100 nm high and ~ 200 nm wide at the base and each contains an ID with a cross section of ~ 10 nm as shown in Fig. 1(b). The IDs were found to originate at the film/substrate interface and extend to the film surface with a constant cross sectional area between 5–20 nm in diameter. The IDBs were along the $\{10\bar{1}0\}$ planes, similar to IDs found in other films grown by MBE^{5,6} and MOCVD.^{2,4}

GaN layers grown by MOCVD with a similar surface morphology have been investigated by Daudin *et al.*² using ion channeling and convergent beam electron diffraction. They found that the IDs associated with pyramids at the surface of their samples were oriented (0001) (or Ga terminated), while the bulk of the matrix was oriented (000 $\bar{1}$) (or

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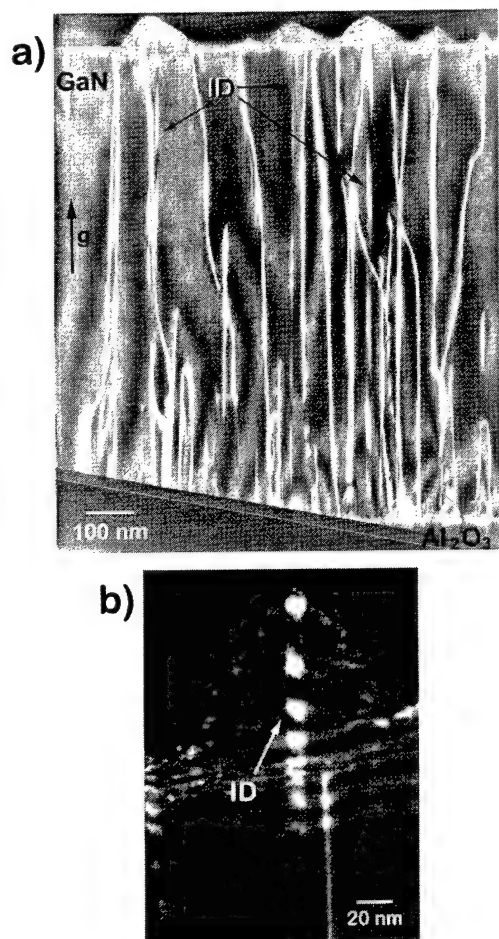


FIG. 1. Multiple dark field images taken with $g=(0002)$ near the $(11\bar{2}0)$ zone axis for a film grown under Ga-rich conditions with atomic hydrogen. (a) Lower magnification of entire film thickness. (b) Higher magnification of pyramid at surface showing the central inversion domain.

N terminated). In contrast, they were also able to grow "flat" samples which were single phase (0001) and contained no IDs. Hillock formation apparently results from the higher growth rate of the ID located at its midpoint. The resultant strain at the boundary may also enhance the growth rate of the opposite phase and dictate the final surface morphology. Perhaps this non-planar growth region is a source of point defects which degrade the properties of samples grown under N-rich conditions.⁷

We believe that our IDs are oriented (0001) in analogy with the MOCVD results.⁷ This is supported by a recent study by Smith *et al.*¹¹ who found by STM measurements for films grown by rf-plasma MBE using similar techniques that the polarity of the matrix was N terminated. If our assessment is correct, however, the Ga-terminated surface can have a significantly higher growth rate than a N-terminated surface, in agreement with the speculations of Middleton *et al.*¹² One implication is that if the bulk matrix is nucleated to be Ga-terminated and a N-terminated ID is of small cross section, the possibility exists for overgrowth of the ID under N-rich conditions, removing this defect after growth of a sufficient layer thickness.

Samples grown under excess Ga were found to be free of pyramidal hillocks, even for films containing high densities of IDBs. An example is given in Fig. 2(a) corresponding to a

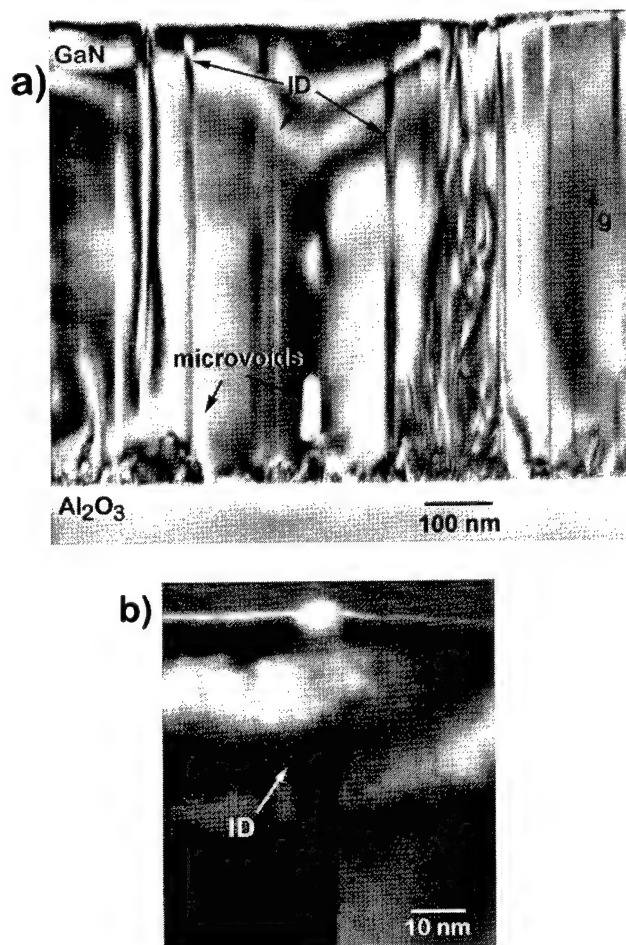


FIG. 2. Multiple dark field images taken with $g=(0002)$ near the $(11\bar{2}0)$ zone axis for a film grown under Ga-rich conditions without atomic hydrogen. (a) Lower magnification of entire film thickness. (b) Higher magnification of an inversion domain at the surface.

sample grown under highly Ga-rich conditions. Again, IDBs were observed to originate at the film/substrate interface and propagate through the growing layer. In sharp contrast to N-rich growth, no significant hillock features were observed at the intersection of the IDB with the surface of the GaN. However, as shown in Fig. 2(b), the surface of the ID is found to be 2 nm higher than the surrounding matrix, indicating a slight enhancement in the growth rate in the region near the ID. This height difference is significantly less than in the film shown in Fig. 1(b). Thus, Ga-rich growth appears to suppress the growth-rate differential between the two different polarities of GaN, directly leading to the observed smoother surface morphologies.

The formation of IDBs is not an intrinsic property of MBE growth of GaN, but is apparently related to nucleation conditions. At least one sample grown in this study had an IDB concentration of $<5\%$, with an ID cross section of <20 nm. As shown here, the IDs initiate within the buffer layer and are localized laterally with a relatively small cross section, indicating a localized nucleation site. We obtain different densities of IDBs with what we believe are identical nucleation conditions. This could be due the presence of steps in the sapphire surface as suggested previously,¹ defects in the substrate surface itself from remnant polishing damage, or possibly related to high energy ions present in the

nitrogen flux itself. (With respect to the latter, we have measured a small but finite flux of nitrogen ions with energies >25 eV from our nitrogen source.¹³)

Nitridation of the sapphire surface prior to growth influences the microstructure and morphology in MOCVD layers.^{14–16} While nitridation is readily accomplished using an ECR plasma source for the active nitrogen,^{17,18} our laboratory and others^{8,19} have not observed any evidence of nitridation with an rf-plasma source for exposures up to one hour. A recent study by Heinlein *et al.* indicate that the nitridation process is very slow with an rf-plasma source, taking up to 5 h for the formation of one monolayer.²⁰ It may be that nitridation is accomplished by ions with sufficient energy for atomic displacement. The slow nitridation rate would thus be due to the small residual high energy ion content found in an rf-plasma source, while the faster nitridation rate observed with ECR-plasma sources naturally comes about from the significantly higher ion density. The seemingly random nucleation of IDs for layers grown by rf-plasma MBE could thus be accounted by inhomogeneous nitridation on the substrate from residual high energy ions during buffer layer nucleation. Experiments are underway in our laboratory to see how ion content affects sapphire nitridation. It may be that by eliminating ions from our rf-plasma source we can suppress ID formation for what we believe is growth of (0001) GaN.

Alternatively, nitridation of the sapphire may allow the subsequent growth of (0001) GaN. Correct choice of which polar surface to grow can have a marked influence on layer properties, as has been observed in other binary semiconductors. At this time, it is not clear which polarity is best for MBE growth.

Other microstructural features can also be observed in the TEM images that depend on the growth conditions. For the sample grown under Ga-rich conditions with atomic H [Fig. 1(a)], a higher density of mixed and screw dislocations were found compared to the sample grown under Ga-rich conditions without atomic H [Fig. 2(a)]. In addition microvoids are present in Fig. 2(a) that are faceted along the $\{10\bar{1}0\}$ and $\{10\bar{1}1\}$ planes. These voids, found to be elongated along the c axis up to 100 nm in length and 20–50 nm wide, may be related to Ga condensation during growth.

In conclusion, the differences in surface morphology observed for different growth conditions could be directly linked to the presence of IDs in our GaN layers. Nitrogen-rich growth and growth under atomic hydrogen enhanced the growth rate of IDs with respect to the surrounding matrix, whereas growth under Ga-rich conditions resulted in a more

nearly equal growth rate. IDs apparently originated in the initial nucleation layer, and were stable with respect to layer growth. A possible cause of the inversion domains is inhomogeneous nitridation on the sapphire substrate due to a remnant high energy ion content in the nitrogen flux from the rf-plasma source.

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¹L. T. Romano, J. E. Northrup, and M. A. O'Keefe, *Appl. Phys. Lett.* **69**, 2394 (1996).

²B. Daudin, J. L. Rouviere, and M. Arley, *Appl. Phys. Lett.* **69**, 2480 (1996).

³X. H. Wu, L. M. Brown, D. Kapolnek, S. Keller, B. Keller, S. P. DenBaars, and J. S. Speck, *J. Appl. Phys.* **80**, 3228 (1996).

⁴Z. Liliental-Weber, H. Sohn, N. Newman, and J. Washburn, *J. Vac. Sci. Technol. B* **13**, 1578 (1995).

⁵L. T. Romano, B. S. Krusor, R. Singh, and T. D. Moustakas, *J. Electron. Mater.* **26**, 285 (1997).

⁶Zhonghai Yu, S. L. Buczowski, N. C. Giles, T. H. Myers, and M. R. Richards-Babb, *Appl. Phys. Lett.* **69**, 2731 (1996); S. L. Buczowski, Zhonghai Yu, M. R. Richards-Babb, N. C. Giles, T. H. Myers, and L. T. Romano, *Mater. Res. Soc. Symp. Proc.* **449**, 197 (1997).

⁷H. Reichert, R. Averbeck, A. Graber, M. Schienle, U. Straub, and H. Tews, *Mater. Res. Soc. Symp. Proc.* **449**, 149 (1997).

⁸R. Beresford, K. S. Stevens, Q. Cui, A. Schwartzman, and H. Cheng, *Mater. Res. Soc. Symp. Proc.* **449**, 361 (1997).

⁹M. Richards-Babb, S. L. Buczowski, Zhonghai Yu, and T. H. Myers, *Mater. Res. Soc. Symp. Proc.* **395**, 237 (1996).

¹⁰D. Lederman, Zhonghai Yu, T. H. Myers, and M. R. Richards-Babb, *Appl. Phys. Lett.* **71**, 368 (1997).

¹¹A. R. Smith, R. M. Feenstra, D. W. Grove, J. Neugebauer, and J. E. Northrup (unpublished).

¹²P. G. Middleton, C. Trager-Cowan, A. Mohammed, K. P. O'Donnell, W. Van Der Stricht, I. Moerman, and P. Demeester, *Mater. Res. Soc. Symp. Proc.* **449**, 471 (1997).

¹³EPI MBE Products Group (St. Paul, MN), Application Note 97-3 (August, 1997).

¹⁴S. Keller, B. P. Keller, Y.-F. Wu, B. Heying, D. Kapolnek, J. S. Speck, U. K. Mishra, and S. P. DenBaars, *Appl. Phys. Lett.* **68**, 1525 (1996).

¹⁵N. Grandjean, J. Massies, and M. Leroux, *Appl. Phys. Lett.* **69**, 2071 (1996).

¹⁶K. Uchida, A. Watanabe, F. Yano, M. Kouguchi, T. Tanaka, and S. Minagawa, *J. Appl. Phys.* **79**, 3487 (1996).

¹⁷T. D. Moustakas and R. J. Molnar, *Mater. Res. Soc. Symp. Proc.* **281**, 753 (1993).

¹⁸M. E. Lin, B. N. Sverdlov, and H. Morkoc, *J. Appl. Phys.* **74**, 5038 (1993).

¹⁹C. Heinlein, J. Grepstad, H. Riechert, and R. Averbeck, *Mater. Sci. Eng. B* **43**, 253 (1997).

²⁰C. Heinlein, J. Grepstad, T. Berge, and H. Riechert, *Appl. Phys. Lett.* **71**, 341 (1997).

The Influence of Growth Conditions, Inversion Domains and Atomic Hydrogen on Growth of (0001) GaN by Molecular Beam Epitaxy

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ABSTRACT

Growth of GaN by rf-plasma molecular beam epitaxy leads to different surface morphologies for nitrogen-stable growth versus gallium-stable growth. Nitrogen-stable growth produces a granular surface morphology with many samples having a significant density of pyramidal hillocks. In contrast, gallium-stable growth results in a flat surface morphology. The hillocks were directly linked to the presence of inversion domains which originated in the nucleation layer. Nitrogen-stable growth and growth under atomic hydrogen enhanced the growth rate of inversion domains with respect to the surrounding matrix, while growth under Ga-stable conditions resulted in a more nearly equal growth rate. Evidence is presented suggesting that hydrogen may stabilize the surface of growing GaN.

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I. Introduction

The potential applications of blue and ultraviolet optoelectronic devices based on GaN have been recognized for many years.[1] Recent advances in epitaxial GaN growth by metal organic chemical vapor deposition (MOCVD) have lead to the first demonstration of a blue laser based on GaN.[2] Rapid progress in this direction is also being accomplished by molecular beam epitaxy (MBE) growth using active nitrogen species.[3,4,5] Several issues remain to be resolved. As noted by Tarsa *et al.* [6], it is becoming clear that certain aspects of growth phenomenon relating to GaN are universal in nature, particularly the predominant effects of the ratio of Ga to nitrogen during growth. For example, growth of GaN under Ga-stable conditions has resulted in improved structural, electrical and optical properties with smoother surface morphologies when compared to N-stable growth[6-10].

Heteroepitaxial GaN layers grown by any technique contain a high dislocation density as well as other classes of defects such as nanopipes, voids and inversion domain boundaries (IDBs). Inversion domains (IDs), consisting of regions of GaN with the opposite polarity to the primary matrix, have been observed to some extent in GaN grown by all techniques [11-17]. However, there is some indication that IDBs are both more common and more stable during growth of GaN by molecular beam epitaxy (MBE) [7,15]. We present the results of a study of IDBs in GaN grown by rf-plasma assisted MBE detailing how different growth conditions determine the effect IDs have on surface morphology.

One of the obvious differences between MOCVD and MBE growth of GaN is the presence of hydrogen, primarily as a component of the molecule supplying the nitrogen. Most studies have either ignored the effect of hydrogen on the growth, or have only considered its effects in compensating p-type dopants such as Mg.[18] Hydrogen may, however, be altering the growth kinetics as well. We also report the effects of the presence of atomic hydrogen on the growth kinetics of GaN. We present evidence that atomic hydrogen can have a significant effect on the growth kinetics of GaN when the growth is limited by the amount of active nitrogen present.

II. Experimental

The GaN layers for this study were grown at West Virginia University by MBE in a custom system. A standard MBE source (EPI-40M) provided the Ga flux. A cryogenically-cooled rf plasma source (Oxford Applied Research CARS-25) operating at either 500 or 600W was used to produce the active nitrogen flux. All layers reported here were grown with a nitrogen flow rate of 6 sccm, controlled by a mass flow controller, resulting in a system background pressure of 6×10^{-5} Torr during growth.

Atomic hydrogen was produced using a thermal cracker (EPI). Typically, 1×10^{-6} Torr beam equivalent pressure (BEP) of hydrogen was passed through the thermal source operating at 9.5 amperes, although the hydrogen flux was varied for several samples. Literature supplied with the source indicated a dissociation efficiency of about 5% for this operating condition. Therefore, the sample was exposed to both atomic and molecular hydrogen during growth.

Determination of substrate temperature and growth rate were important in this study. A spring-loaded Type K thermocouple was in intimate contact with the back of the molybdenum sample block with the substrate mounted with indium. Prior to this sequence of growths, an additional thermocouple was attached to the front of a mounted substrate and a calibration curve determined between the front and back thermocouples. This calibration was routinely checked by using the melting points of various metals, oxide desorption from GaAs, and the use of an optical pyrometer for the higher temperatures. Temperature determination was reproducible to ± 5 °C.

Relative measurements of sample growth rates were performed by measuring interference effects in reflectance from the growing layer using 680 nm light from a semiconductor laser. These measurements were converted to an absolute growth rate by using total sample thickness and the growth time. The thickness at the sample's center was determined from interference fringes in optical transmittance measured using a Cary-14 spectrophotometer. This determination of total thickness was found to agree with values determined by transmission electron diffraction (TEM).

All samples were grown on c-plane sapphire substrates (Union Carbide Crystal Products). Prior to growth, the substrates were degreased and etched in a

phosphoric/sulfuric (1:3) acid mixture heated to 80°C. Based on our earlier study,⁹ buffer layers were grown by heating the substrate to 730 °C under an atomic hydrogen flux for 20 minutes and then cooling to 630 °C for the growth of a 200 Å thick GaN buffer layer under a Ga flux of 5.0×10^{-7} Torr (BEP) with a 6 sccm nitrogen flow at 600 W. Buffer layer growth was initiated by simultaneous exposure to the Ga and N flux. This nucleation layer was then annealed at 730 °C for 20 minutes under nitrogen flux, cooled to the growth temperature, and growth was resumed. These conditions represent buffer layer growth under highly Ga-stable conditions. However, after the 730 °C anneal, examination of the buffer layers by AFM indicated continuous films with no evidence of Ga condensation. This procedure led exclusively to the nucleation and growth of (0001)-oriented (or N-terminated) GaN as determined using the polarity-indicating etch described by Seelmann-Eggebert *et al.* [19] As discussed in a later section, this determination is consistent with other observations.

Atomic force microscopy (AFM) and TEM were used to study surface morphology and its relation to microstructure in a series of GaN layers grown by MBE on (0001) sapphire. Detailed descriptions of other layer characterizations are given elsewhere. [7,20,21] Cross section TEM (XTEM) studies were performed with a JEOL 3010 microscope operated at 300kV on samples that were prepared by polishing and then ion milling to electron transparency. TEM images were taken under various diffraction conditions including multiple dark field imaging with $g = \pm (0002)$ along either the $\langle 11\bar{2}0 \rangle$ or $\langle 10\bar{1}0 \rangle$ axis in order to reveal inversion domains [11]. AFM was performed in air using a Digital Instruments Nanoscope II.

III. Growth Conditions and Surface Morphology

As reported by our group[7] as well as others [8,9,3,6], growth under Ga-stabilized conditions vs. N-stabilized conditions leads to a drastically different surface morphology. When varying the Ga flux with all other growth conditions fixed, two different regimes are indicated by RHEED, as shown in Fig. 1. For a higher Ga-flux, a

RHEED pattern was observed consisting of well-defined, truncated streaks similar to Fig. 1(a). Such a pattern indicates a surface that is locally smooth on an atomic scale, and is indicative of two-dimensional (2-D) growth. Lowering the Ga flux lead yielded a transition to a spot pattern such as shown by Fig. 1(b). This transition to a pattern indicative of three-dimensional (3-D) growth implies a surface that is rough on an atomic scale. For a short duration of 3-D growth, the transition was reversible and 2-D growth could be recovered by increasing the Ga flux. The transition was fairly abrupt, occurring over a small change in Ga flux, about 0.5×10^{-7} Torr beam equivalent pressure (BEP).

Measurement of the GaN growth rate as a function of Ga flux indicates that the RHEED pattern change can be attributed to a transition from Ga-stable to N-stable growth. In Figure 2, the growth rate is plotted as a function of Ga beam equivalent pressure. The nitrogen flow rate was held constant at 6 sccm, 600W. For an active nitrogen flux in excess or equal to the Ga flux, one would expect a linear increase in growth rate with increasing Ga flux since any excess nitrogen will desorb. When the surface ratio of Ga to active nitrogen is increased beyond unity, then either Ga condensation will occur if the desorption rate for excess Ga is low or the growth rate will become fairly constant with increasing Ga flux if the excess Ga desorption rate is high. A linear increase in growth rate was observed in going between 1.3 to close to 4.0×10^{-7} Torr BEP Ga. A least-squares fit to these points, including a fixed point of zero growth at zero flux, is represented by the solid line in Figure 1. The growth rate did not increase significantly above this Ga-flux, indicating that the growth was now limited by the amount of active nitrogen available. As indicated in the figure, we actually observed a decreasing growth rate for increasing Ga flux, similar to that reported by Held *et al.* [10] for GaN growth with ammonia-based MBE. This region of constant (or decreasing) growth rate represents Ga-stable growth, whereas the linearly increasing region represents the nitrogen-stable case. A series of experiments where the Ga-flux was varied and growth rate measured *in-situ* indicated that the 2-D to 3-D transition and plateau in growth rate occurred at the same Ga-flux, to within 0.5×10^{-7} Torr BEP. For the conditions shown in Fig. 2, this occurred for a Ga flux around 5×10^{-7} Torr BEP.

As we reported earlier, both AFM [7] and x-ray surface scattering [21] measurements indicate a very smooth surface morphology for GaN grown under Ga-stable conditions, with an rms. surface roughness less than 1 nm. Features less than 1 nm in height are difficult to resolve with our AFM. In contrast, samples grown under N-stable conditions exhibited a highly textured surface morphology, with rms. surface roughness of 10 to 20 nm. Such samples displayed a coarse surface morphology dominated by grain-like features as indicated by Fig. 3. Smaller Ga-to-N flux ratios appeared to produce a finer-scale morphology. On many samples, a high density of triangular shaped pyramidal hillocks was also observed. As discussed in Section IV, the hillock features were associated with inversion domains.

A recent study by Tarsa *et al.* [6] produced essentially identical results for homoepitaxial growth of GaN on MOCVD-grown GaN using similar growth conditions. Of interest here is that the MOCVD-GaN used by Tarsa *et al.* is believed to have a (0001)-oriented (or Ga-terminated) growth surface, whereas ours is (000 $\bar{1}$). The similarity in morphology and growth modes indicate that the fundamental mechanisms governing growth kinetics may be independent of orientation. Our results, coupled with that of Tarsa *et al.*, suggest that N-stable conditions result in low surface adatom mobility resulting in "quenched" growth and leading to statistical roughening of the surface.[22] In contrast, Ga-stable conditions appears to promote surface adatom mobility, resulting in a larger surface diffusion length and leading to two-dimensional growth.

IV. Inversion Domains and Surface Morphology

Samples grown either under N-stable conditions or under an atomic hydrogen flux often exhibit a highly textured surface morphology consisting of pyramidal hillocks. Figure 4 is an AFM micrograph illustrative of such a surface completely dominated by the triangular hillocks, while Figure 5 is from a sample exhibiting isolated hillocks. The hillocks have a triangular cross-section that range from 0.2 to 0.5 μm on a side for 1 μm thick samples, and between 150-250 nm in height. Increasing the Ga/N ratio (without

atomic H) results in smaller size pyramids. These features were not observed for highly Ga-rich conditions.

Figure 6 shows a XTEM image of a hillock from a 1.0 μm thick sample grown under an atomic hydrogen flux with a surface that contained a high density of pyramid-shaped hillocks. The hillocks were ~ 100 nm high and ~ 200 nm wide at the base and each contained an ID with a cross-section of ~ 10 nm as shown in Fig. 6. The IDs were found to originate at the film/substrate interface and extend to the film surface with a constant cross sectional area between 5-20 nm. The IDs were along the $\{10\bar{1}0\}$ planes, similar to IDs found in other films grown by MBE [15,7] and MOCVD [12,14]. What is amazing is the pronounced effect the small ID has on surface morphology. In terms of sample volume, the ID density in the sample shown in Figure 4 is less than 5%, yet effects associated with the IDs totally dominate the sample's morphology.

GaN layers grown by MOCVD with a similar surface morphology have been investigated by Daudin *et al.* [12] using ion channeling and convergent beam electron diffraction. They found that the IDs associated with pyramids at the surface of their samples were oriented (0001) (or Ga- terminated), while the bulk of the matrix was oriented (000 $\bar{1}$) (or N-terminated). In contrast, they were also able to grow "flat" samples which were single phase (0001) and contained no IDs. Hillock formation apparently results from the higher growth rate of the ID located at its midpoint. The resultant strain at the boundary may also enhance the growth rate of the opposite phase and dictate the final surface morphology. Perhaps this non-planar growth region is a source of point defects which degrade the properties of samples grown under N-stable conditions. [8].

We believe our IDs are oriented (0001) similar to the MOCVD results [12]. This is supported by a recent study by Smith *et al.* [23] who found by STM measurements for films grown by rf-plasma MBE using similar techniques that the polarity of the matrix was N-terminated. If our assessment is correct, however, the Ga-terminated surface can have a significantly higher growth rate than a N-terminated surface, in agreement with the speculations of Middleton *et al.* [24]. The growth rate differential is also consistent with recent results on MOCVD GaN reported by Liliental-Weber *et al.*, [26] where (000 $\bar{1}$)-oriented IDs grow at a slower rate than the (0001)-oriented matrix, leading to oriented

"pits" that could be described as inverted hillocks. One implication is that if the bulk matrix is nucleated to be Ga-terminated and a N-terminated ID is of small cross-section, the possibility exists for overgrowth of the ID under N-stable conditions, removing this defect after growth of a sufficient layer thickness. Thus, the observation that inversion domains are stable during MBE growth of GaN may be due to the relative differential growth rate on (0001)-oriented material.

Samples grown under excess Ga were found to be free of pyramidal hillocks, even for films containing high densities of IDBs. Again, IDBs were observed to originate at the film/substrate interface and propagate through the growing layer. In sharp contrast to N-stable growth, no significant hillock features were observed at the intersection of the IDB with the surface of the GaN. However, as reported earlier [16] the surface of the ID is found to be 2 nm higher than the surrounding matrix, indicating a slight enhancement in the growth rate in the region near the ID. This height difference is significantly less than in the film shown in Fig. 6. Thus, Ga-stable growth appears to suppress the growth-rate differential between the two different polarities of GaN, directly leading to the observed smoother surface morphologies.

The formation of IDBs is not an intrinsic property of MBE growth of GaN, but is apparently related to nucleation conditions. We have grown several samples under N-stable conditions that only exhibit the granular texture indicated by Fig. 3, without the presence of hillocks. The IDs initiate within the buffer layer and are localized laterally with a relatively small cross-section, indicating a localized nucleation site. We obtain different densities of IDBs with what we believe are identical nucleation conditions. This could be due the presence of steps in the sapphire surface as suggested previously [11], defects in the substrate surface itself from remnant polishing damage, or possibly related to high energy ions present in the nitrogen flux itself. (With respect to the latter, we have measured a small but finite flux of nitrogen ions with energies > 25 eV from our nitrogen source[25])

VI. Influence of Atomic Hydrogen

The presence of atomic hydrogen dramatically increases the growth rate of GaN grown under Ga-stable conditions, as shown in Figure 7. The growth rates indicated by the solid triangles in Figure 7 were grown under a total (atomic and molecular) hydrogen flux of 1×10^{-6} Torr BEP. The other growth parameters remained the same as for the corresponding filled-circle case. Note that the growth rate was essentially tripled for the most Ga-rich growth. In contrast, growth under nitrogen-stable conditions did not exhibit an enhanced growth rate. A recent study by Daudin and Widmann [26] indicate a similar increase for the growth rate of AlN grown under Al-stable conditions due to the introduction of hydrogen.

The increase in growth rate for the Ga-stable cases was not very sensitive to the overall hydrogen flux. Changing the hydrogen flux from 0.5 to 2.0×10^{-6} Torr BEP gave the same value for the increase in the growth rate. To see if the increased growth rate originated with molecular hydrogen, samples were grown under hydrogen flux with the cracker turned off. The resulting samples exhibited identical growth rates to the GaN grown without hydrogen, indicating that molecular hydrogen is not significantly affecting the growth kinetics. Also, to see if the atomic hydrogen was possibly forming active species with molecular nitrogen, an attempt was made to grow under an atomic hydrogen flux with the rf power turned off on the nitrogen source. The resulting GaN growth rate, if non-zero, was too small to be detected.

The increase in growth rate for Ga-stable conditions is apparently related to the presence of atomic hydrogen. The growth rate enhancement itself is not sensitive to the actual atomic hydrogen overpressure within the limits we investigated, but depends only on its presence. It is well known that hydrogen easily bonds to the surface of other semiconductor systems, such as silicon[27] or diamond.[28] Also, there is evidence that atomic hydrogen alters the growth kinetics in GaAs.[29] One possibility is that the atomic hydrogen becomes loosely bonded to the growing GaN surface. Nitrogen atoms adsorbed on the surface are then attracted by this hydrogen layer, resulting in an increased nitrogen residence time. The longer residence time increases the probability that a Ga

atom will diffuse to within an interaction distance of the nitrogen, and thus enhance the growth rate of GaN. Thus, the atomic hydrogen could be increasing the effective active nitrogen concentration. Of interest, the surface morphology for samples grown under atomic hydrogen more nearly resembled N-stable growth. We also observed the same enhanced ID growth rate as for N-stable growth, resulting in the pyramidal structures. The morphology of the background was generally granular, similar to Fig. 3. Surface morphology became smoother only for the most Ga-stable-plus-hydrogen conditions, where rms. roughness values of 1 to 2 nm were measured. These observations, along with the increased growth rate, is consistent with shifting the growth kinetics towards a more nitrogen-stable growth.

The effect of hydrogen on growth can be illustrated another way. Many groups using MBE have reported a trend similar to that shown by the filled triangles in Fig. 8. That is, the growth rate of GaN is fairly constant for fixed flux values as the temperature is increased up to some value, typically in the range 700 to 800 °C, where the growth rate rapidly decreases. This temperature depends on the absolute magnitudes of the flux and may be related to the onset of rapid desorption of Ga from the growing surface. [30] The growth rate can be recovered for temperatures up to the limit of our heater by introducing atomic hydrogen, as indicated by filled circles in Fig. 8. Thus, the atomic hydrogen may be affecting the desorption kinetics of Ga as well. The stabilizing effect of atomic hydrogen for the growing GaN surface deserves additional study.

V. Conclusion

In conclusion, our nucleation and growth conditions produce (0001)-oriented GaN on basal-plane sapphire. For this orientation, N-stable growth produced a granular, 3-D surface while Ga-stable growth resulted in a smoother surface morphology indicative of 2-D growth. The presence of pyramidal hillocks could be directly linked to the presence of IDs in our GaN layers. Nitrogen-stable growth and growth under atomic hydrogen enhanced the growth rate of IDs with respect to the surrounding matrix, whereas growth under Ga-stable conditions resulted in a more nearly equal growth rate. IDs apparently

originated in the initial nucleation layer, and were stable with respect to layer growth due to a differential growth rate. We have demonstrated that the presence of atomic hydrogen can have a significant effect on the growth rate of GaN under Ga-stable conditions.

Hydrogen may stabilize the growing GaN surface at higher temperatures

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References

1. See, for example, J. Pankove, *Mater. Res. Soc. Symp. Proc.* **97**, 409 (1987).
2. S. Nakamura, M. Senoh, S. Nagahama, N. Iwasa, T. Yamada, T. Matsushita, H. Kiyoku and Y. Sugimoto, *Jap. J. Appl. Phys.* **35**, L74 (1996).
3. T. D. Moustakas, T. Lei, and R. J. Molnar, *Physica B* **185**, 36 (1993).
4. M. Smith, G.D. Chen, J.Z. Li, J.Y. Lin, H.X. Jiang, A. Salvador, W.K. Kim, O. Aktas, A. Botchkarev and H. Morkoç, *Appl. Phys. Lett.* **67**, 3387 (1995).
5. M.A.L. Johnson, Zhonghai Yu, C. Boney, W.C. Hughes, J.W. Cook, Jr, J.F. Schetzina, H. Zhao, B.J. Skromme, and J.A. Edmond, *Mat. Res. Symp. Proc.* **449**, 215 (1997).
6. E.J. Tarsa, B. Heying, X.H. Wu, P. Fini, S.P. DenBaars and J.S. Speck, *J. Appl. Phys.* **82**, 5472 (1997).
7. Zhonghai Yu, S.L. Buczkowski, N.C. Giles, T.H. Myers and M.R. Richards-Babb, *Appl. Phys. Lett.* **69**, 2731 (1996); S.L. Buczkowski, Zhonghai Yu, M.R. Richards-Babb, N.C. Giles, T.H. Myers and L.T. Romano, *Mat. Res. Soc. Symp.* **449**, 197 (1997).
8. H. Reichert, R. Averbeck, A. Graber, M. Schienle, U. Straub, and H. Tews, *Mater. Res. Soc. Symp. Proc.* **449**, 149 (1997).
9. R. Beresford, K.S. Stevens, Q. Cui, A. Schwartzman, and H. Cheng, *Mater. Res. Soc. Symp. Proc.* **449**, 361 (1997).
10. R. Held, D.E. Crawford, A. M. Johnston, A. M. Dabiran, and P.I. Cohen, *J. Electron. Mater.* **26**, 272 (1997).
11. L.T. Romano, J. E. Northrup, and M. A. O'Keefe, *Appl. Phys. Lett.* **69**, 2394 (1996).
12. B. Daudin, J. L. Rouviere, and M. Arley, *Appl. Phys. Lett.* **69**, 2480 (1996).
13. X. H. Wu, L. M. Brown, D. Kapolonek, S. Keller, B. Keller, S. P. DenBaars, and J. S. Speck, *J. Appl. Phys.* **80**, 3228 (1996).
14. Z. Liliental-Weber, H. Sohn, N. Newman, and J. Washburn, *J. Vac. Sci. Technol. B* **13**, 1578 (1995).
15. L.T. Romano, B. S. Krusor, R. Singh, and T. D. Moustakas, *J. Electron. Mater.* **26**, 285 (1997).

16. L.T. Romano and T.H. Myers, Appl. Phys. Lett. **71**, 3486 (1997).
17. V. Potin, P. Ruterana and G. Nouet, J. Appl. Phys. **82**, 2176 (1997).
18. M.S. Brandt, J.W. Ager, III, W. Gotz, N.M. Johnson, J.S. Harris, Jr, R.J. Molnar and T.D. Moustakas, Phys. Rev. **B 49**, 14758 (1994).
19. M. Seelmann-Eggebert, J.L. Weyher, H. Obloh, H. Zimmermann, A. Rar and S. Porowski, Appl. Phys. Lett. **71**, 2635 (1997).
20. M. Richards-Babb, S.L. Buczkowski, Zhonghai Yu, And T.H. Myers, Mater. Res. Soc. Symp. Proc. **395**, 237 (1996).
21. D. Lederman, Zhonghai Yu, T.H. Myers and M.R. Richards-Babb, Appl. Phys. Lett. **71**, 368 (1997).
22. D.L. Smith in Thin Film Deposition (McGraw Hill, New York, 1995), pp. 119-184.
23. A. R. Smith, R. M. Feenstra, D. W. Greve, J. Neugebauer, and J. E. Northrup, Phys. Rev. Lett. **79**, 3934 (1997).
24. P.G. Middleton, C. Trager-Cowan, A. Mohammed, K.P. O'Donnell, W. Van Der Stricht, I. Moerman and P. Demeeser, Mat. Res. Symp. Proc. **449**, 471 (1997)
25. EPI MBE Products Group (St. Paul, MN), Application Note 97-3 (August, 1997).
26. Z. Liliental-Weber, Y. Chen, S. Ruvimov and J. Washburn, Phys. Rev. Lett. **79**, 2835 (1997)
27. See, for example, V.A. Burrows, Y.J. Chabal, G.S. Higashi, K. Raghavachari and S.B. Christman, Appl. Phys. Lett. **53**, 998 (1988).
28. See, for example, *The Properties of Natural and Synthetic Diamond*, edited by J.E. Field (Academic Press, London, 1992) and references therein.
29. H. Shimomura, Y. Okada and M. Kawabe, Jpn. J. Appl. Phys. **31**, L628 (1992).
30. C.R. Jones, T. Lei, R. Kaspi and K.R. Evans, Proceedings of the 1995 Fall MRS Meeting, Symposium AAA, (unpublished).

Figure Captions

Figure 1. RHEED along $[2\bar{1}10]$ of GaN grown under (a) Ga-stable conditions (b) N-stable conditions.

Figure 2. Growth rate of GaN for various values of Ga flux..

Figure 3. AFM micrograph of GaN grown under N-stable conditions. Surface features had an rms. surface roughness of about 30 nm.

Figure 4. AFM micrograph of a GaN surface dominated by triangular hillocks. The pyramidal structures were about 100 nm high.

Figure 5. AFM micrograph of a GaN surface showing isolated hillocks. The hillocks were about 120 nm high. This sample was grown slightly Ga-stable, resulting in a smooth (rms. roughness about 1 nm) between the hillocks.

Figure 6. Dark field TEM image of a pyramid at the surface taken with $g = (0002)$ near the $(11\bar{2}0)$ zone axis of a film grown under Ga-stable conditions with atomic hydrogen showing the central inversion domain.

Figure 7. Effect of atomic hydrogen on the growth rate of GaN for various conditions.

Figure 8. Influence of atomic hydrogen on the temperature-dependence of the growth of GaN.

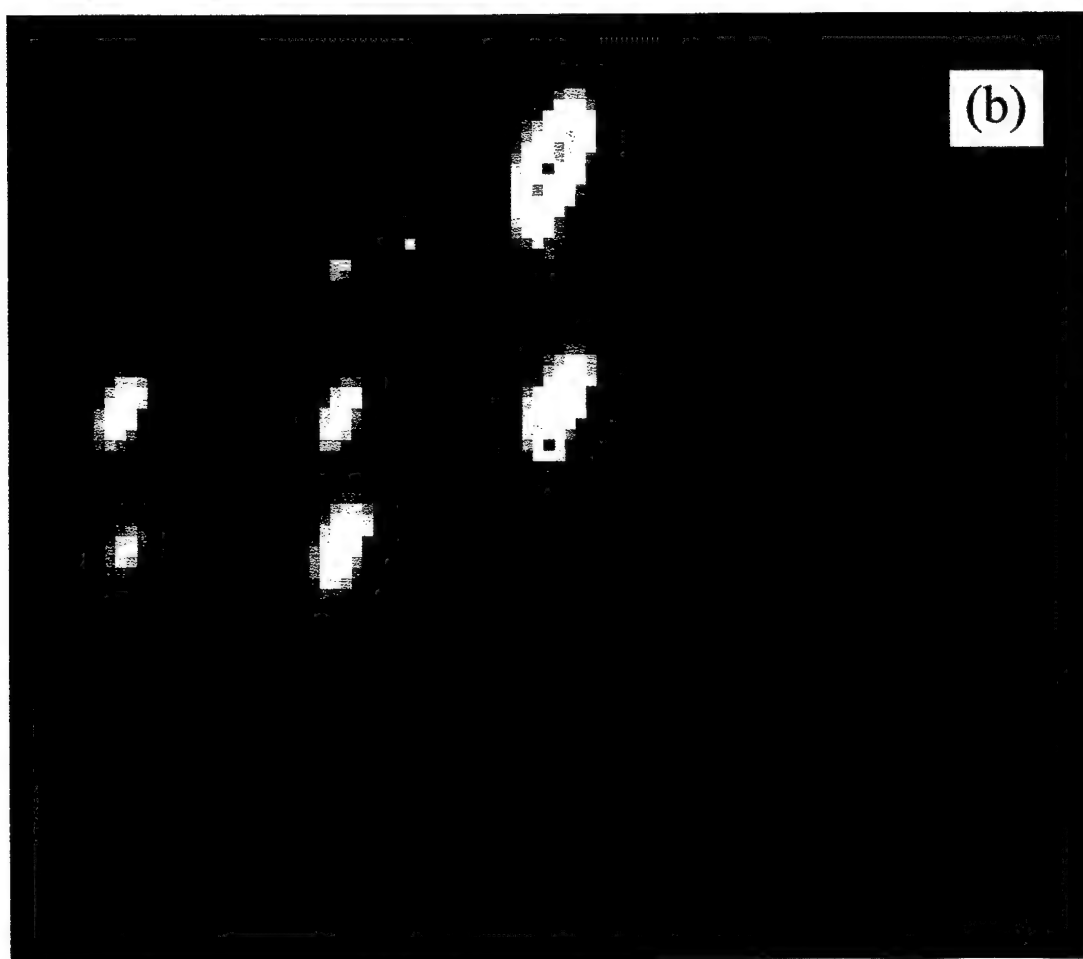
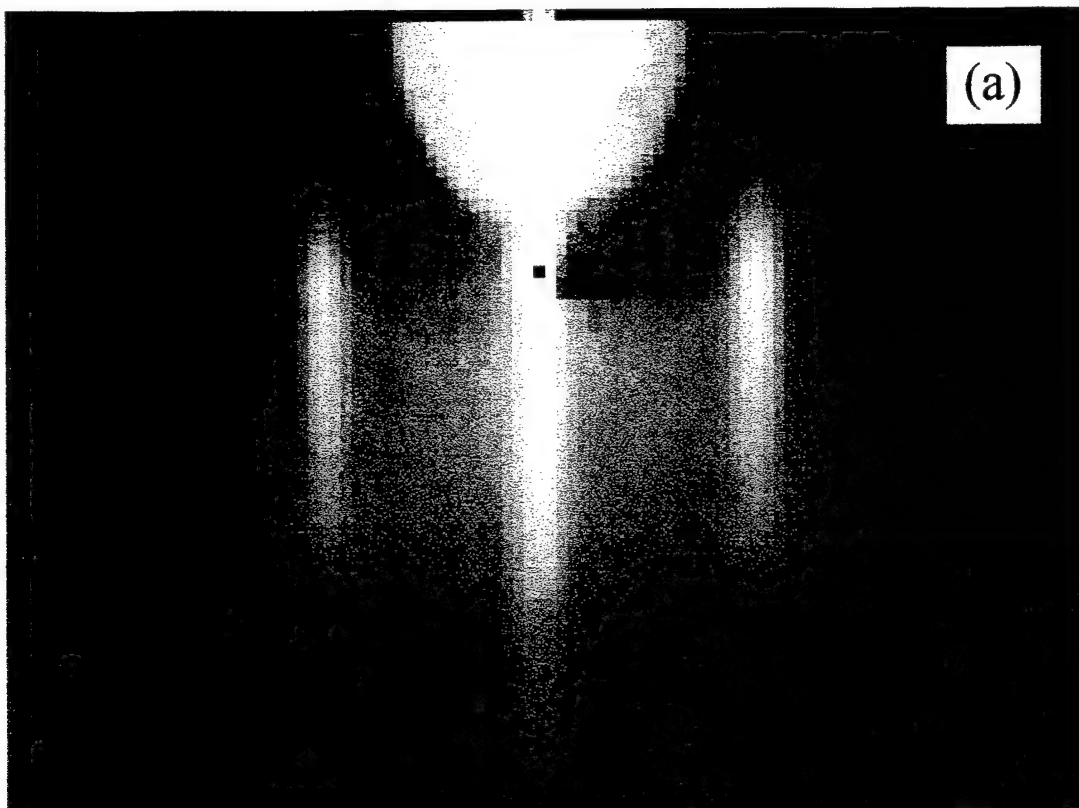


Figure 1

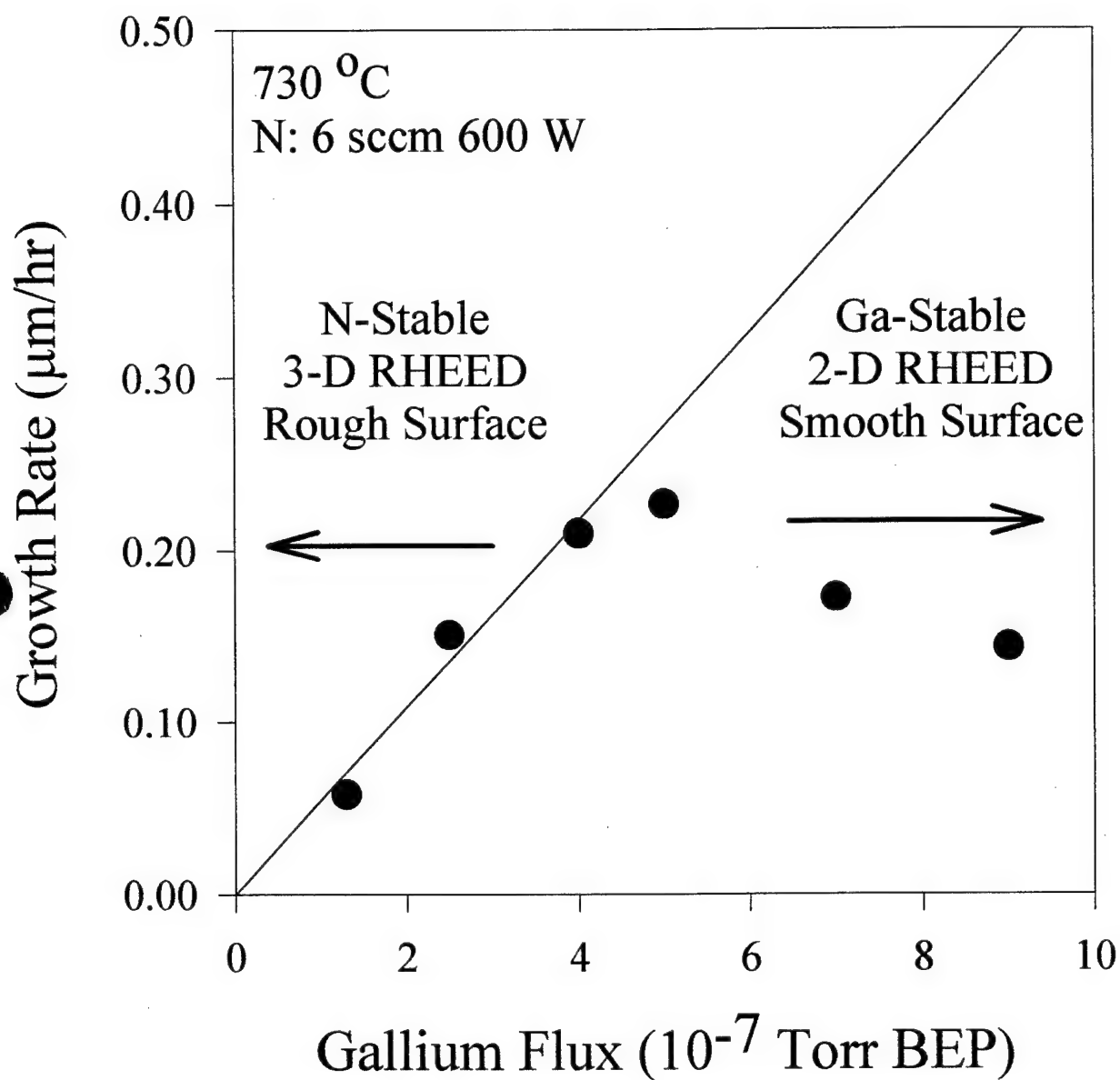


Figure 2

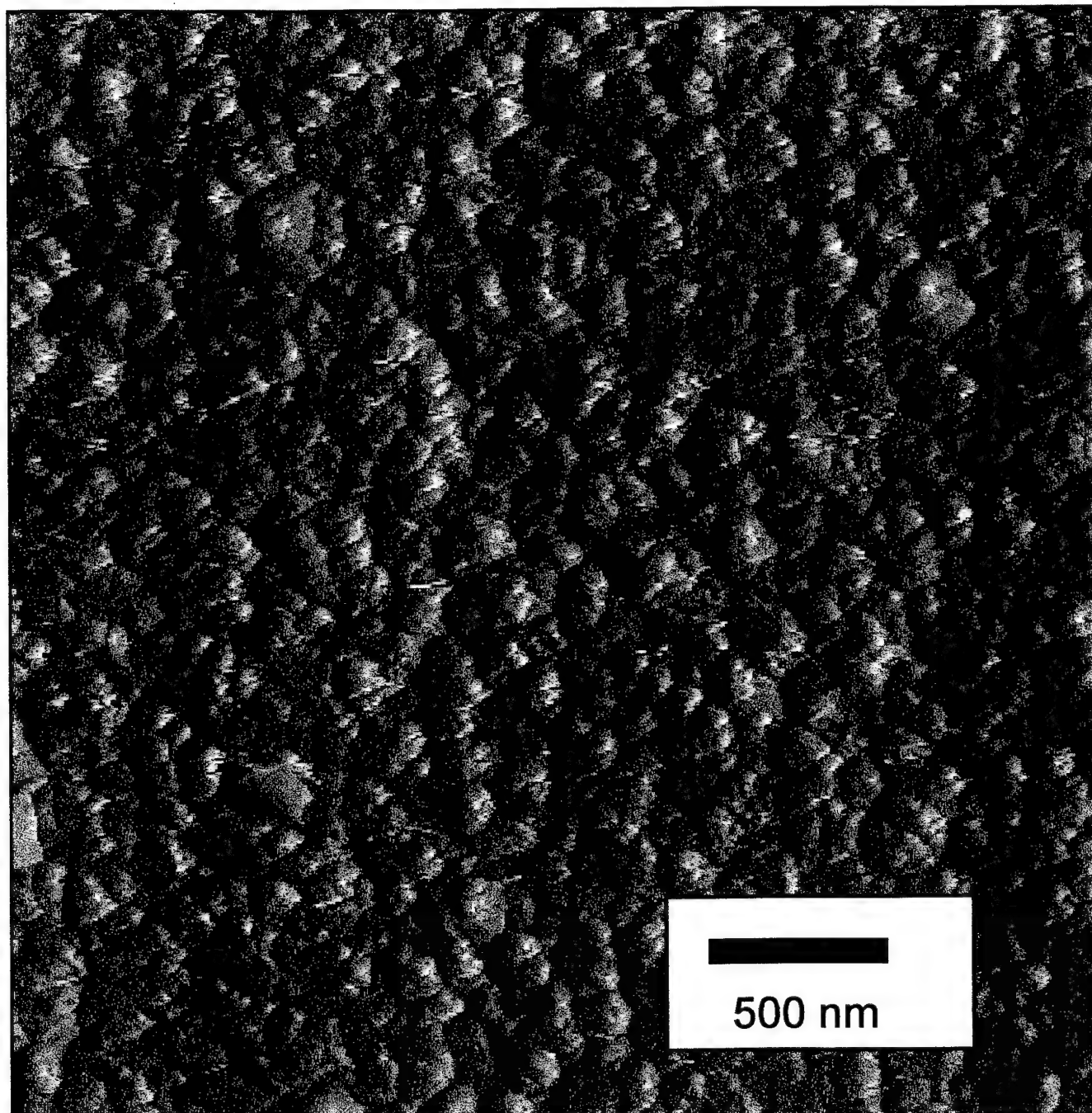


Figure 3.

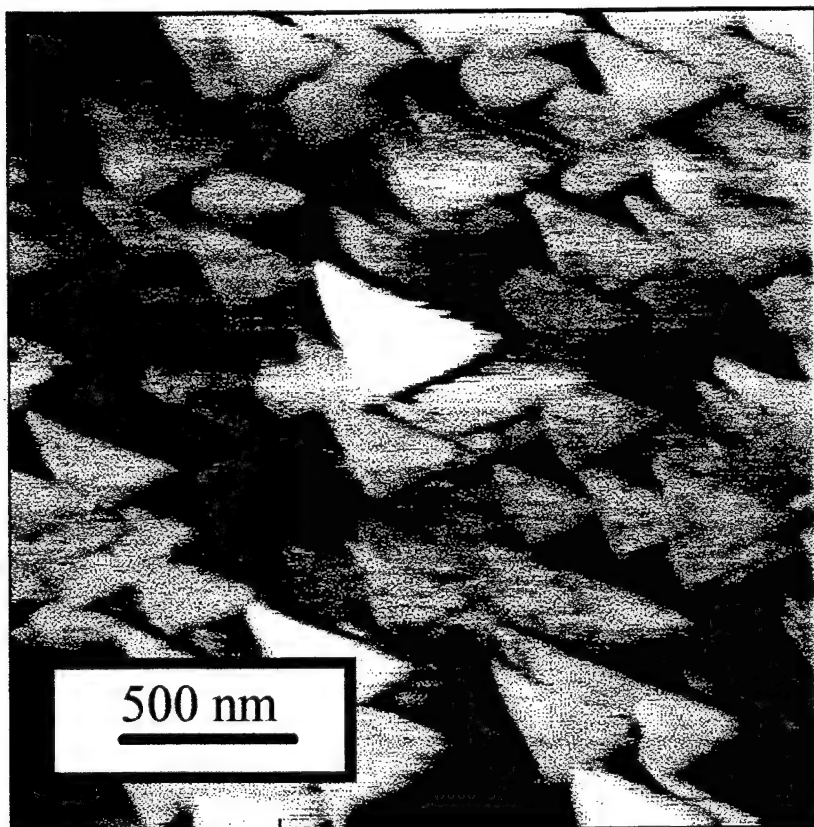


Figure 4

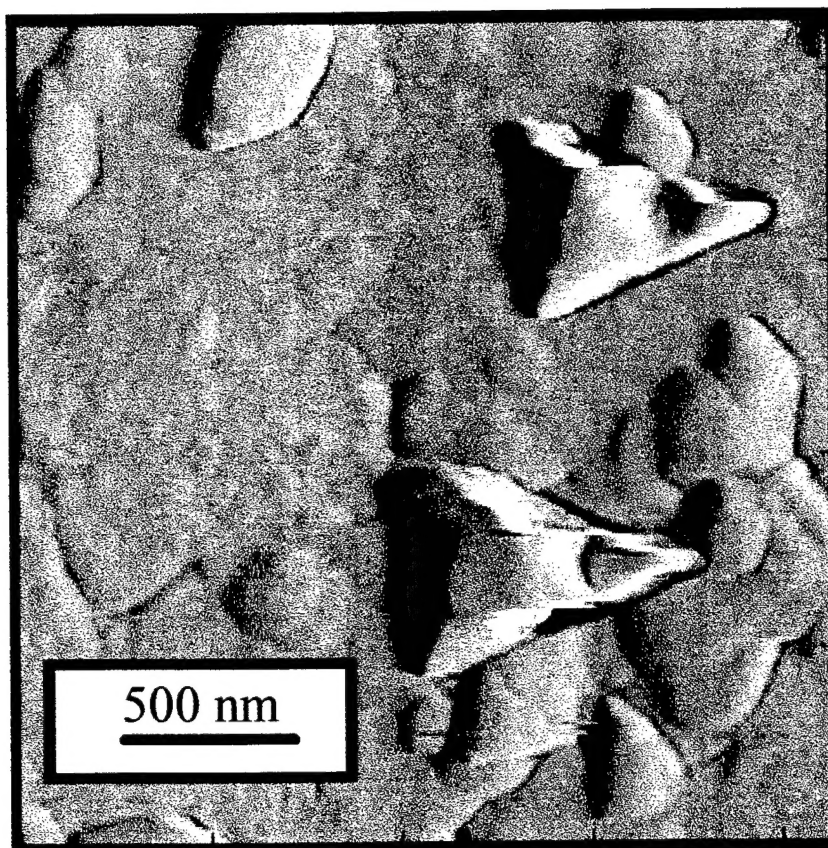


Figure 5

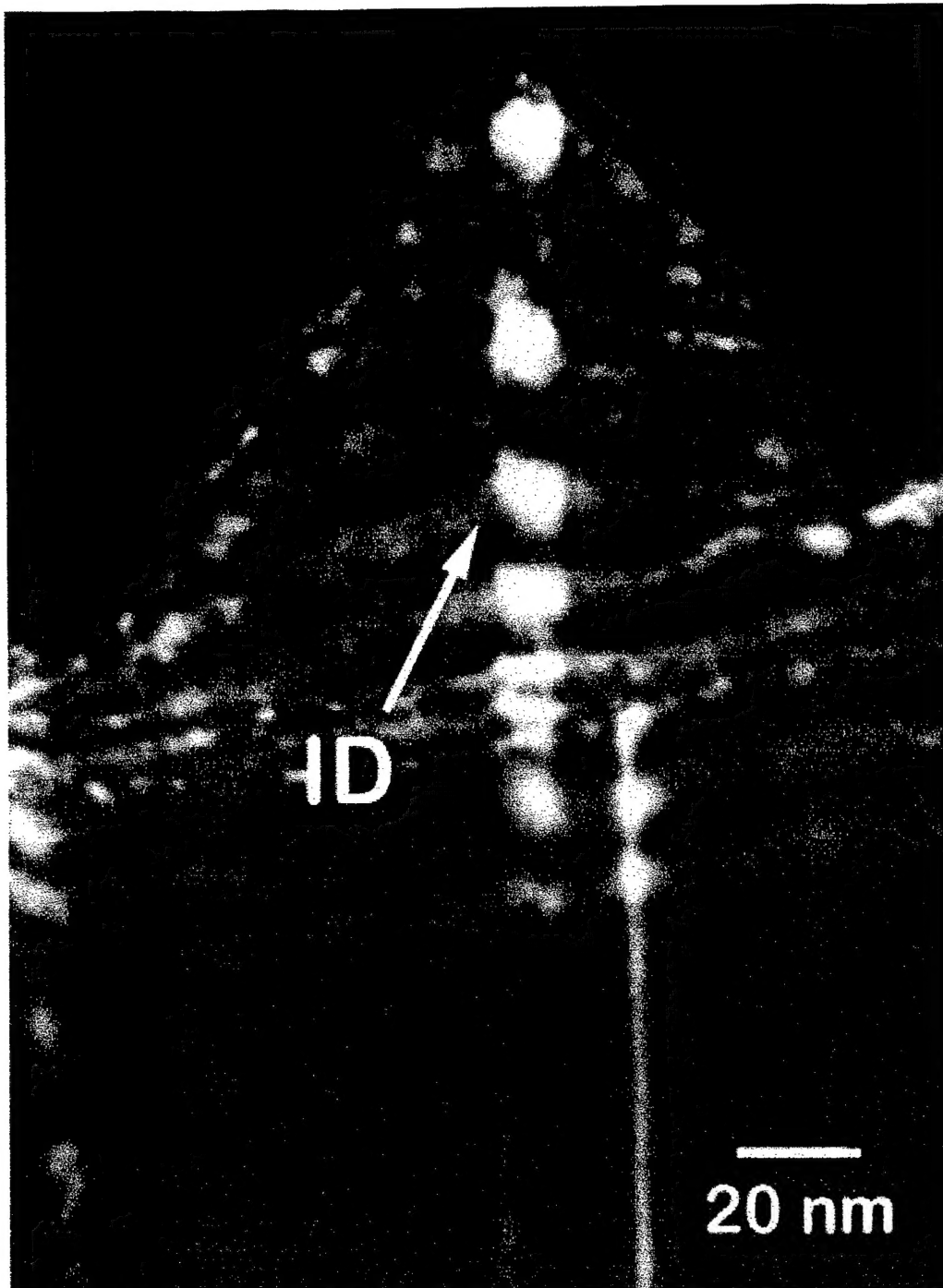


Figure 6

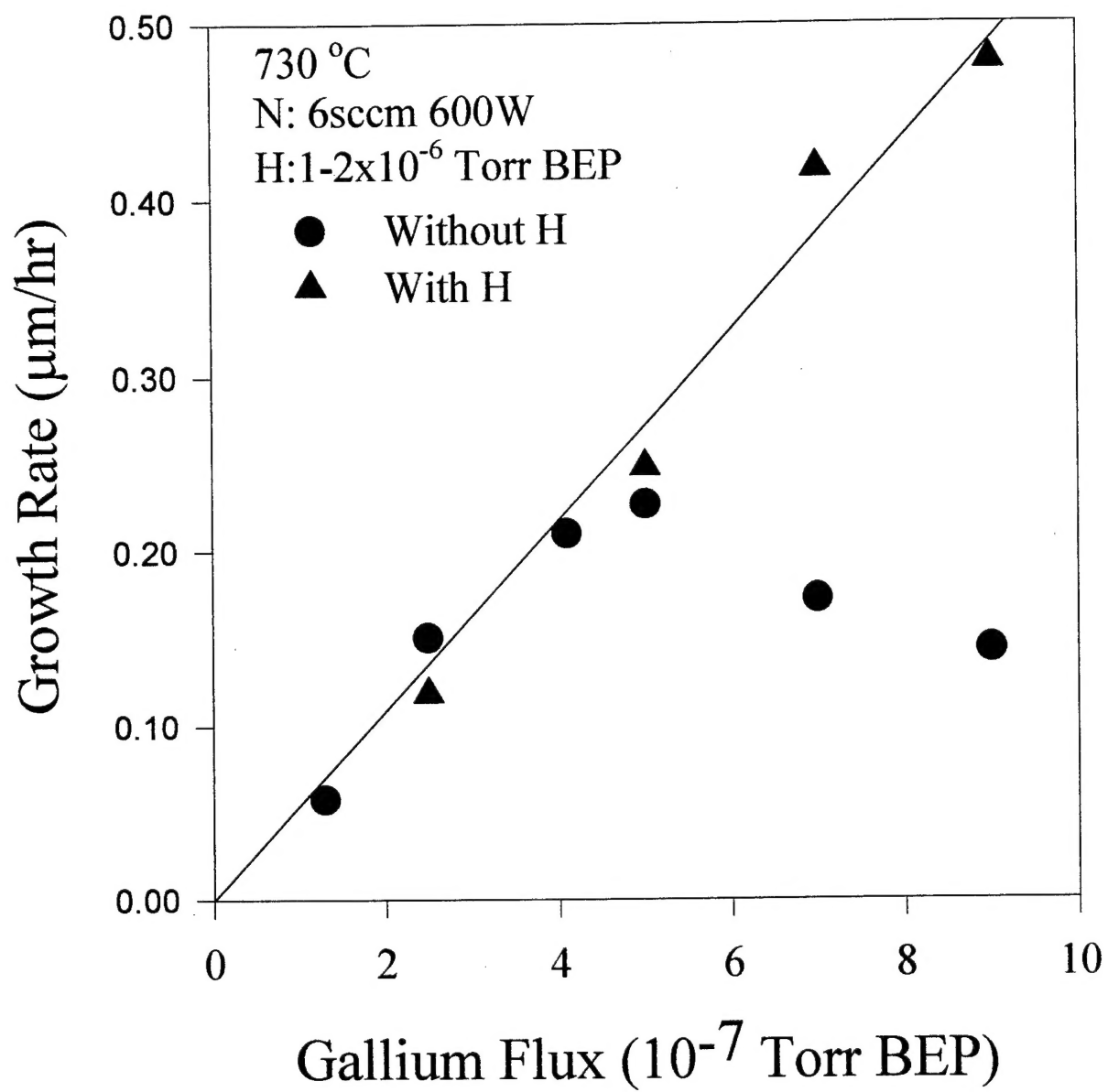


Figure 7

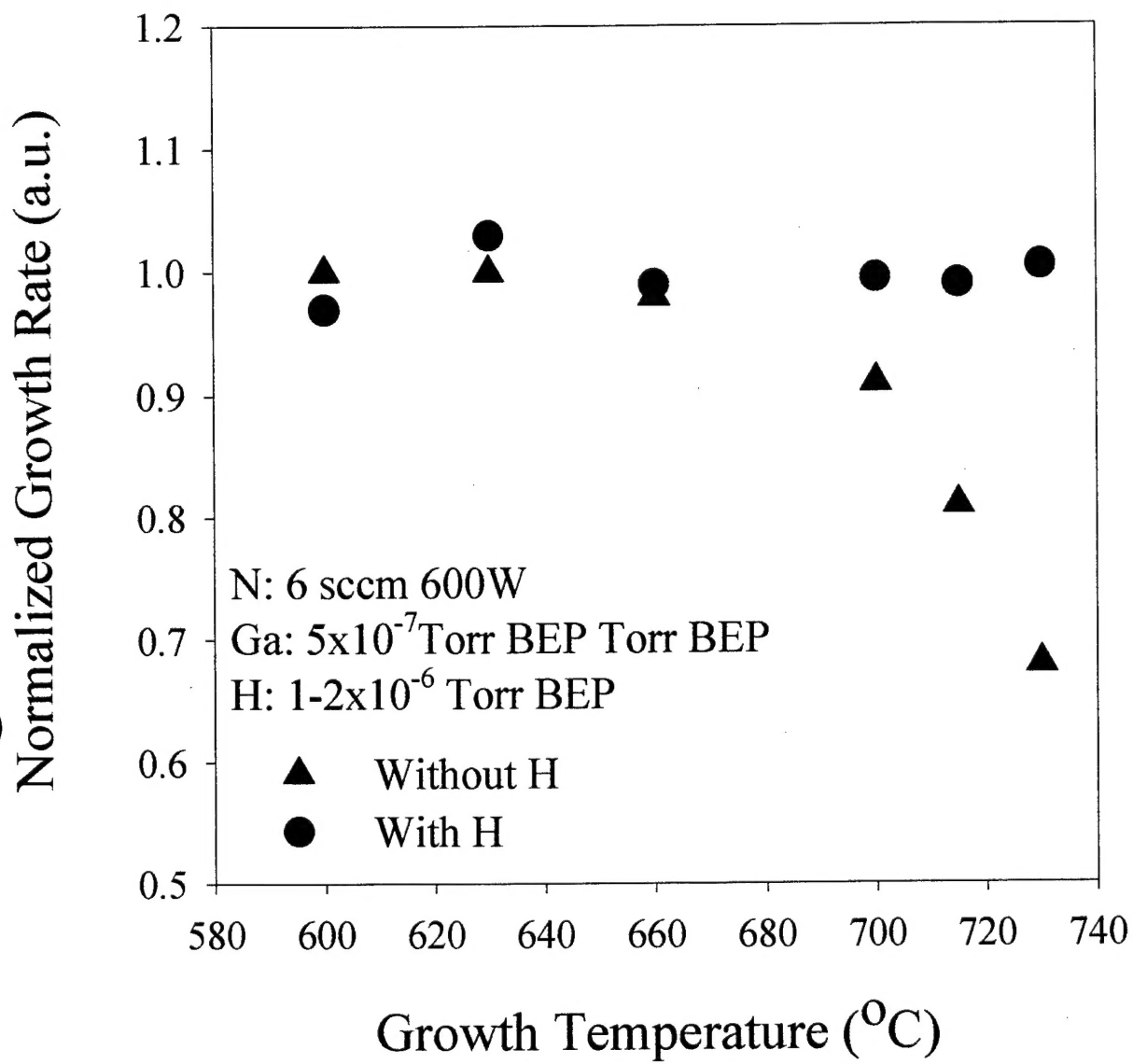


Figure 8